A TEXT-BOOK OF
INORGANIC CHEMISTRY.
VOLUME IV.

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## A TEXT-BOOK OF INORGANIC CHEMISTRY.

J. NEWTON FRIEND, D.Sc., F.I.C., Ph.D., Carnegue Gold Medalllat

An Introduction to Modern Inorganic Chemistry. By J. NLWION FRIEND, D.Sc. (Blum), Ph.D. (Wurz), H. F. V. LITTIE, B.Sc. (Lond.), A.R.C.S., Chief Chemist to Thorium, Ltd., W. E. S. TURNER, D.Sc. (Lond.). VOLUME I. Third Edition Pp 1-xv 385 12s. 6d PART II. The Inert Gases (Group 0 in the Periodic Table). By H. V. A. Briscot, D Sc (Lond.), A R C S The Alkali Metals and their Congeners (Group I of the VOLUME II In the Press Periodic Table) By A. Jamiesos Walker, Ph.D. (Heid.). The Alkaline Earth Metals and their Associates (Group VOLUME III. II of the Periodic Table) By Miss M S LESLIE, D.Sc. Aluminium and its Congeners, including the Rare Earth Metals (Group III. of the Periodic Table) By VOLUME IV. Second Edition. Pp. i-xx + 485. H. F. V LITTLE, B.Sc. (Lond.), A R.C.S., Chief Chemist to Thornum, Ltd. Carbon and its Allies (Group IV of the Periodic Table). VOLUME V By R. M CAVEN, D.Sc (Lond), F.I.C Second Impression. Pp. i-xxi + 468VOLUME VI. Nitrogen and its Congeners (Group V. of the Periodic Table) By E B R. PRIDEAUX, M.A., D.Sc. PART I. Oxygen. By J. Newton Friend, D.Sc., and Douglas F. Twiss, D.Sc., F.I.C.

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## THE PERIODIC TABLE.

PERIODS.	GROUP O.	GROUP I.	GROUP IL	GROUP I	rla -				
		н	-	- GROUP II	GROUP IV.	GROUP V.	GROUP VI.	GROUP VII.	GROUP VIII.
First short period	He 4.00	L <sub>1</sub> 6 94	G	B 11 B	. 1	N	0		
Second short period .	Ne 20 2	Na 23 00	Mg 24 3	g Al	Si	14·01	16 00	19 0	
First Even series .	A 39 39	K 59 10	Ca	32 27 ·:	T <sub>1</sub>	31 04	32 06	Cl 35·46	
period (Odd ,, .		Cu 63 57	40.07 Zn 65.3	44.1 Ga	48·1 Ge	51 0 As	Cr 52 0 S <sub>"</sub>	Mn 54 93 Br	Fe Co Ni 55 84 58 97 58 6
Second   Even series . long   Oilu ,	Kr 52 9	Rb 85 45	Sr 87.63	Y 88.7	Zr 90 6	74 96 Cb	79 2 Mo	79 92	
Thud (Even series	!	107 88	Cd 112	In	Sn	93·1 Sb 120 2	96·0	4 /	Ru Rh Pd 101:7 102:9 106:7
longs   Even series	X 130 2	Cs 132 81	Ba 137 37					126 92	
Fourth Even series .	·			THE RARE EARTH METAL		Ta			•
period (Odd ,, .	.	Au 197 2	Нg 200 б	7,	Pb 207·20	181 5 Bı	184 0	.	Os Ir Pt 190·9 193·1 195·2
Fifth long period .	Nt 222 4		Ra 226.0		Th	205 0		-	
formulæ of oxides formulæ of hydrides		R <sub>2</sub> O RH	RO RH,	R <sub>2</sub> O <sub>3</sub>	RO,	R <sub>3</sub> O <sub>5</sub>	RO,		
Volume in this series of text-books.	1			RH <sub>3</sub>	RH4	RH,	RH <sub>2</sub>	R <sub>2</sub> O <sub>7</sub> RH	RO <sub>4</sub>
		2	3	4	5	opted in this Tal	7	e 8	9

The International Atomic Weights for 1947 are adopted in this Table.

# A TEXT-BOOK OF INORGANIC CHEMISTRY.

J. NEWTON FRIEND, D.Sc., Ph D., F.I.C.,

## VOLUME IV.

ALUMINIUM AND ITS CONGENERS, including

THE RARE EARTH METALS

BY

H. F. V. LITTLE, B Sc (LOND), A R.C S., D.I C. (CHEF CHI MIST TO THORICM, LTD.)

With Frontispicce, 2 Plates, and 44 3llustrations in the Text

SECOND EDITION





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1921

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#### GENERAL INTRODUCTION TO THE SERIES.

DURING the past few years the civilised world has begun to realise the advantages accruing to scientific research, with the result that an ever-in-reasing amount of time and thought is being devoted to various branches of science

No study has progressed more rapidly than chemistry. This science may be divided roughly into several branches namely, Organic, Physical, Inorganic, and Analytical Chemistry. It is impossible to write any single text-book which shall contain within its two covers a thorough treatment of any one of these branches, owing to the vast amount of information that has been accumulated. The need is rather for a series of text-books dealing more or less comprehensively with each branch of chemistry. This has already been attempted by enterprising firms, so far as physical and analytical chemistry are concerned, and the present series is designed to meet the needs of imaganic chemists. One great advantage of this procedure lies in the fact that our knowledge of the different sections of science does not progress at the same rate. Consequently, as soon as any particular part advances out of proportion to others, the volume dealing with that section may be easily revised or rewritten as occasion requires.

Some method of classifying the elements for treatment in this way is clearly essential, and we have adopted the Periodic Classification with slight alterations, devoting a whole volume to the consideration of the elements in each vertical column, as will be evident from a glance at the scheme in the Frontispiece.

In the first volume, in addition to a detailed account of the Elements of Group 0, the general principles of Inorganic Chemistry are discussed. Particular pains have been taken in the selection of material for this volume, and an attempt has been made to present to the reader a clear account of the principles upon which our knowledge of modern Inorganic Chemistry is based.

At the outset it may be well to explain that it was not intended to write a complete text-book of Physical Chemistry. Numerous excellent works have already been devoted to this subject, and a volume on such lines would scarcely serve as a suitable introduction to this series. Whilst Physical Chemistry deals with the general principles applied to all branches of theoretical chemistry, our aim has been to emphasise their application to Inorganic Chemistry, with which branch of the subject this series of text-books is exclusively concerned. To this end practically all the illustrations to the laws and principles discussed in Volume 1. deal with inorganic substances.

Again, there are many subjects, such as the methods employed in the accurate determination of atomic weights, which are not generally regarded

as forming part of Physical Chemistry. Yet these are subjects of supreme importance to the student of Inorganic Chemistry, and are accordingly included in the Introduction.

. Hydrogen and the ammonium salts are dealt with in Volume II, along with the Elements of Group I The position of the rare earth metals in the Periodic Classification has for many years been a source of difficulty. They have all been included in Volume IV., along with the Elements of Group III., as this was found to be the most suitable place for them.

Many alloys and compounds have an equal claim to be considered in two or more volumes of this series, but this would entail unnecessary duplication. For example, alloys of copper and tin might be dealt with in Volumes II. and V. respectively. Similarly, certain double salts—such, for example, as ferrous ammonium sulphate—might very logically be included in Volume II under ammonium, and in Volume IX under iron. As a general rule this difficulty has been overcome by treating complex substances, containing two or more metals or bases, in that volume dealing with the metal or base which belongs to the highest group of the Periodic Table—For example, the alloys of copper and tin are detailed in Volume V. along with tin, since copper occurs earlier, namely in Volume II.—Similarly, feirous animonium sulphate is discussed in Volume IX. under iron, and not under ammonium in Volume II.—The ferrocyandes are likewise dealt with in Volume IX.

But even with this arrangement it has not always been found easy to adopt a perfectly logical line of treatment. For example, in the chromates and permanganates the chromium and manganese function as part of the acid radicles and are analogous to sulphur and chlorine in sulphates and perchlorates; so that they should be treated in the volume dealing with the metal acting as base, namely, in the case of potassium permanganate, under potassium in Volume II—But the alkali permanganates possess such close analogies with one another that separate treatment of these salts hardly seems desirable. They are therefore considered in Volume VIII.

Numerous other little irregularities of a like nature occur, but it is hoped that, by means of carefully compiled indexes and frequent cross-referencing in the texts of the separate volumes, the student will experience no difficulty in finding the information he requires

Particular care has been taken with the sections dealing with the atomic weights of the elements in question. The figures given are not necessarily those to be found in the original memoirs, but have been recalculated, except where otherwise stated, using the following fundamental values —

```
    Hydrogen = 1:00762,
    Oxygen = 16:000.

    Sodium = 22:996.
    Sulphur = 32:065.

    Potassium = 39:100.
    Fluorine = 19:015.

    Silver = 107:880
    Chlorine = 35:457.

    Carbon = 12:003
    Bromme = 79:916.

    Nitrogen = 14:008.
    Jodine = 126:920.
```

By adopting this method it is easy to compare directly the results of earlier investigators with those of more recent date, and moreover it renders the data for the different elements strictly comparable throughout the whole series.

Our aim has not been to make the volumes absolutely exhaustive, as this would render them unnecessarily bulky and expensive; rather has it been to contribute concise and suggestive accounts of the various topics,

and to append numerous references to the leading works and memoirs dealing with the same. Every effort has been made to render these references accurate and reliable, and it is hoped that they will prove a useful feature of the series. The more important abbreviations, which are substantially the same as those adopted by the Chemical Society, are detailed in the subjoined list.

In order that the series shall attain the maximum utility, it is necessary to arrange for a certain amount of uniformity throughout, and this involves the suppression of the personality of the individual author to a corresponding extent for the sake of the common welfaire. It is at once my duty and my pleasure to express my sincere appreciation of the kind and ready mainer in which the authors have accommodated themselves to this task, which, inthout their hearty co-operation, could never have been successful. Finally, I wish to acknowledge the unfailing courtesy of the publishers, Messrs Charles Griffin & Co., who have done everything in their power to render the work straightforward and easy.

J NEWTON FRIEND

March 1917

#### PREFACE.

The present volume aims at giving a tolerably complete and readable a bougt of the morganic chemistry of the elements of the third group of the Periodic Table, together with cerium and the tervalent rare earth elements. Where they are of particular interest or importance, however, various organic compounds of these elements have also been described, particularly in dealing with the rare earth elements.

So far as was possible without overburdening the text, the results of modern physico-chemical work have been included. It is hoped that the bibliography relating to the pure chemistry—morganic and physico-chemical—of the elements described is practically complete and that no important references are omitted. The discussions of mineralogical, technical, and analytical chemistry are of necessity very brief, but it is hoped that the bibliographies attached to them will be of service to the reader.

The chapters dealing with the rare earth elements together constitute more than half the book, and form the most complete account of the chemistry of these elements that has yet appeared in English. No apology is needed for the somewhat lengthy discussion of the spectra of these elements. The nature of the cathodic phosphorescence spectra has been purposely discussed in considerable detail, and it is hoped that the description of

discussed in considerable detail, and it is hoped that the description of Urbain's beautiful researches on this subject may assist Eughsh chemists in realising that the atmosphere of mystery and romance which they have so long associated with these elements has at last been dispelled

In the preparation of the text, the Dictionaries of Morley and Muir, Watts, Wurtz, and Thorpe and the Handbooks of Abegg, Danmer, Fehling, Gmelin, Ladenburg, and Mossan have been freely consulted. The Abstracts published by the English and American Chemical Societies have also been of great service. A very considerable proportion of the text, however, has been prepared directly from the original memons. Books consulted for information on special topics are usually mentioned in toothotes, in addition to those, Browning's Introduction to the Rayer Elements, Bohm's Die Darstellung der seltenen Erden, and Truchot's Les Terres Rayes may be mentioned. Most of the references cited have been checked by reference to the original journals, etc.

The Author desires to express his deep indebtedness to Dr A. B Searle, the well-known authority on clays, for his kindness in critically revising the proof-sheets of Chapters IV and V., and to thank Professor C. James, of New Hampshire College, U.S.A., Dr M W Travers, F.R.S., and several other gentlemen for preate information on a variety of points. For the preparation of the name index the Author is indebted to Dr J. Newton Friend.

The photographs from which the plates have been prepared were kindly

taken for the Author by Mr K. Hickman, the necessary facilities having been granted by Professor, A. Fowler, F.R.S., of the Imperial College of Science and Technology, and the rare earths having been presented by Professor C. James

Corrections, omissions. or criticisms will be the control of the control

Corrections, omissions, or criticisms will be thankfully received by the Author, who would also be grateful for reprints of original papers bearing on the subjects included in the book

H F. V. LITTLE

Thorium, Lad., Illord Essex, March 1917

Reprinted 1921.

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#### LANTHANUM Properties of the Metal-Alloys

Compounds of Lanthanum—General Characteristics—Thermochemistry
Hydride—Fluoride—Cilioride—Oxychloride—Brounde—Lodide—Perchloride—Enough — Perchloride—Enough — Perchloride—Persulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Nitride—Aride—Nitride—Aride—Nitride—Nitride—Nitride—Aride—Nitride—Carbide—Carbide—Thioganate—Phosphide—Arides—Ariente—Arienate—Carbide—Carbide—Thioganate—Platinocyanide—Acetate—Ethylsulphide—Acetate—Oxalae—Oxalae—Organic Salts—Metabolate

#### PRASEODYMIUM-Properties of the Metal

COMPOUNDS OF PRASFODYMIUM - General Characteristics-Thermochemistry

Hydride—Fluoride—Chlorde—Oxychlorde—Bromde—Iodide—Bromate
—Oxide—Hydroxide—Hydroxide—Oxyde—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Tungstate
—Silicotungstate—Nitride—Nitride—Double Nitrates—Carbide—Carbonate
Ethylsulphide—Acetylacetonate—Oxalate—Organic Sults

#### NEODYMIUM Properties of the Metal

COMPOUNDS OF NEODYMH M - General Characteristics - Thermochemistry

Hydride - Fluoride - Chloride - Oxychloride - Bronnde - Iodide - Bronnde 
Oxide - Hydroxide - Hydrated Perovide - Sulphide - Sulphate - Molybdate

- Tringstate - Silicotingstate - Uranate - Nitride - Nitrate - Double Nitrates

- Carbide - Carbonate - Ethylsulphate - Acetylacetonate - Oxalate - Organic Sules

#### SAMARIUM-Properties of the Metal

Compounds of Samarum—General Characteristics—Thermochemistry.

Fluoride—Chloride—Oxychloride—Subchloride—Bromide—Todide—Subnodide—Bromate—Todate—Periodate—Oxide—Hydroxide—Peroxide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Nitrate—Phosphates—Vanadates—Carbide—Carbonate—Platmocyanide—Thiocyanate—Ethylsulphide—Acetylacetonate—Oxalate—Organic Salts—Orthoborate

#### CHAPTER XIII. The Rare Earth Elements (continued)

THE TERBIUM GROUP.

Salts

EUROPIUM - Chloride - Subchloride - Oxide - Sulphate - Ethylsulphate -Organic salts

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GADOLINIUM—Fluoride—Chloride — Oxychloride — Bromide—Oxide — Hydroxide —Sulphide—Sulphate—Selenite—Selenate—Silcotungstate — Nitrate — Vanadate—Carbonate—Platinocyanide—Oxalate — Ethylsulphate—Acetate —Organic Salts—Metaborate,	
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HOLMIUM—Oxide.	•
YTTRIUM.	
COMPOUNDS OF YTTRIUM—Fluoride—Chloride—Bromde—Chlorate—Perchlorate—Bromate—Iodate—Periodate—Oxide—Hydroxide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Sulphide—Dribhonate—Selenate—Tungslate—Silicotangstate—Nitrate—Phosphates—Carbide—Carbonate—Platinocyanide—Thiocyanide—Ethylsulphide—Acetylacetonate—Organic Salts—Silicate	
$\begin{tabular}{ll} \bf ERBIUM-Oxide-Sulphate-SelemteTungstateSiheotungstateNitratePlatinocyanideOxidateOrganic Salts \\ \end{tabular}$	
THULIUM — Chloride — Bromate — Oxide — Sulphate — Nitrate — Oxalate — Acetylacetonate — Organic Salts.	
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P. 69. Line 7 in § on Aluminum Iodule, for "exclusive" read "explosive." P. 80, 2 ,, ,, Selemde, ,, "sulphate" ,, "selemate" P. 92. ,, 12 from top, ,, "disulphate" ,, "disulphate" P. 95. ,, 10 ,, bottom, ,, "of" ,, "or."	

### LIST OF CHIEF ABBREVIATIONS EMPLOYED. IN THE REFERENCES.

#### AIRTEVIALID TILLI

#### JOT RVAL

	to a set Lemma Lemma
Inter Chem J	American Chemical Johnnal.
Amer J Ser	American Johnnal of Science and Arts
tunt les que i	Anale de la Socied id E-pañola Fisica y Quinnea
$Analy \cdot l$	The Analyst
.Analon	Annalen der Chemie,
Ing Chem	Annales de Chimie (1719-1815, and 1911+)
Ann Chan, and	Annales de Chame analytique appliquée à l'Industrie, a
	1 Agriculture a la Pharmacie, et a la biologie
Ina Chine applicata	Annali di Chinica aigdicata.
Inn Chem Phys	Annales de Chima et de Physique (1816-1913)
'nu l'alset.	Annales des Falsiteatrens
Ann II n	Annales des Minos
Ann Puns .	Annales de Physique (1914+)
	Amaden der Physik
Ann Physik	Archy der Pharmizie
Arch Pharm	Archives des Sciences physiques et naturelles.
Arch Ser phas nat	Arkiy for Kenn. Mineralogi och Geologi
Arkie Kem Meg Ged	Atti della Reale Aceadenna Lancer
Atta R Accord   Line   .	Bereichte der deutschen chemischen Gesellschaft.
Ber .	n to to the transfer of the bank to Cocollectual
Le deut physikal Ger	Berahte der deutschen plasskalischen Gesellschuft
Lall, Acad my P 'q .	Academic royale de l'elgique -Bulletin de la Classe des
	Scheller
Ball Acad See Cracow	Bulletin international de l'Academie des Seiences de Ciacovie.
Bull A ad Ser Petronal	Bulletin de l'Academie Imperiale des Sciences de Petrograd
Intll Something	Bulletin de la Société chimique de France
Rall Soc chine Bela	Bulletin de la Société chimique de Belgique
Ball Son d'En p l'Ind	Bulletin de la Société d'Encour gement pour l'Industrie
nationale	nationale
Bull Soc fram Min.	Bulletin de la Société francaise de Minéralogie
Bull U S. Ged Surrey	Paller and the United States Geological Sulvey
Centr Min .	Centrarblatt f.n. Mineralogie, Geologie, und Paleentologie.
Cham Ind	Die chemische Industrie.
Chem. News	Chemical News
Chem. Wee' blad	Cheni ch Weckblad
Chem. In the drive	Chemiker Zeitung
Chem Zeit .	Ch. m. ches Zentralblatt
Chem Zenti	Comptes rendus hebdomadaires des Séances de l'Académie
Compt rend .	des Seiences
	Dingler's polytechnisches Joannal
Dinge poly I	Gaz, etta chunca italiani.
Huzetla .	Geological Magazine
Geol. Mag.	International Zeitschrift für Metallographie,
Intern. Zedoch Melallegrapher	Nenes Jahrbuch for Mineralegie Geologie, und Palæontologie.
Jahrb Men	
Jahrb Radmaktiv I'e'trond	Jahresberreht über die Fortschritte der Chemie.
Jahr esber	James Berre II and the Forest Champal Society
J Amer. Chem. So	Journal of the American Chemical Society.
J Chim. phys	Journal de Chime physique
J. Coll Sci. Tokyo	Journal of the College of Science, Tokyo
J. Franklin Inst . •	Journal of the Franklin Institute.
J Ind Eng Chem.	Journal of Industrial and Engineering Chemistry.
J. Pharm Chim	Journal de Pharmacie et de Chimie.
J Physique .	Journal de Physique
J. Physical Chem.	Journal of Physical Chemistry.
a anguina	XIX

#### ABBREVIALED TITLE

#### JOURNAL

J. prakt. Chem	Journal fur praktische Chenne.
J. Roy. Agric. Soc	Journal of the Royal Agricultural Society.
J. Russ. Phys Chem Noc.	Journal of the Physical and Chemical Society of Russia.
J. Foc. Chem. Ind .	Journal of the Society of Chemical Industry
J. Sor. Dyers .	Journal of the Society of Dyers and Colourists
J. Washington Acad. Ser	Journal of the Washington Academy of Sciences
Kolloud-Zeitsch	Kolloud-Zeitschrift
K Svenska Vet -Akad Handl.	Kongl Svenska Vetenskaps-Akadennens Handlingar
Mem. Acad Ron. Belg	Mémoires de l'Academie Royale de Belgique
Met. (hem Eng .	Metallurgical and Chemical Engineering
Min. May.	Mineralogical Magazine and Journal of the Mineralogical
activity and the second	Society
Monatsh,	Monatshelte for Chemic and verwandte Theile anderer
	Wissenschaften
Nova Acta Soc. Upsala	Nova Acta Regie Societatis Scientiarum Upsaliensis
Pharm. Zentr. h.	Pharmaceutische Zentralballe
Phil. Mag	Philosophical Magazine (The London, Edinburgh, and
	Dublin)
Phil Trans	Philosophical Transactions of the Royal Society of London
Phys. Review	Physical Review.
Physikal. Zeitsch .	Physikalische Zeitschrift
Pagg Annalen .	Poggendorff's Annalen der Physik und Chemie
Proc. Chem Soc	Proceedings of the Chemical Society
Proc. Amer. Acril	Proceedings of the American Academy of Arts and Sciences.
Proc. Camb Phil Soc.	Proceedings of the Cambridge Philosophical Society.
Proc K Akad Wetensch	Koninklijke Akademie van Wetenschappen te Anisterdam,
Amsterdam	Proceedings (English Version).
Proc Manchester Lit Phil	Proceedings of the Manchester Literary and Philosophical
Soc.	Society
Proc. Roy Suc	Proceedings of the Royal Society of London
Proc Roy. Soc. Lain	Proceedings of the Royal Society of Edinburgh.
Rec trav chrm	Recenil des travanx chimiques des Pays-Bas et de la Belgique.
Schneigyei's J.	Schweigger's Journal Im Chemie und Physik Continued as
	Journal for praktische Cheune)
Ser, Proc Roy Dubl Son	Scientific Proceedings of the Royal Dublin Society
Sitzungsber K. Akud Wiss	Sitzungsberichte der Koniglich-Preussischen Akademie der
Berlin	Wissenschaften zu Berlin
Sitzungshi K Akad. Wiss	Sitzungsberichte der Kaiserhehen Akademie der Wissen-
Wien	schaften, Wien
South African J Sci .	South African Journal of Science
Tech. Papers Bur Stand	Technical Papers, Bureau of Standards, Washington
Tomind, Zeit.	Tonindustric Zeitung
Trans. Amer. Cer Suc	Transactions of the American Ceramic Society.
Trans. Amer Electrochem Soc	Transactions of the American Electrochemical Society.
Trans. Chem Suc	Transactions of the Chemical Society
Trans. Eng. (er. Noc	Transactions of the English Ceramic Society
Trans, Faraday Suc	Transactions of the Faraday Society
Trans Inst Mng. Enq .	Transactions of the Institute of Vining Engineers.
Tach, Min. Mill	Tschermak's Mineralogische Mitteilungen Wiedensam's Annalogische Physik und Chauta
Wred. Annulen	Wiedemann's Annalen der Physik und Chemie Zeitschrift für analytische Chemie
Zertsch anal. Chem .	Zeitschrift für angewandte Chemie
Zertsch angew Chem .	Zertschrift für anorganische Chemie
Zeitsch, anorg. Chem . Zeitsch, Chem. Ind. Kolloule	Zeitschrift für Cheime und Industrie des Kolloide (con-
Benette, Shem. Int. Autoute	timed as Kolloid Zeitschrift)
Zeitsch. Elektrochem	Zeitschrift für Elektrocheime
Zeitsch. Kryst Man.	Zeitschrift im Krystallogiaphie und Mineralogie
Zeitsch. physikal. Chem.	Zertschrift fin physikalische Chemie, Stochionietrie und
zowacie. przystrut. Oneit.	Verwandtschaftslehre.
Zeitsch. wiss. Photochem	Zeitschrift für wissenschaftliche Photographie, Photophysik
201100-11 (60001 2 100000101101	and Photochemie

### TABLE OF DATES OF ISSUE OF JOURNALS.

For the sake of easy reference, a list is appended of the more important journals in chronological order, giving the dates of issue of their corresponding series and volumes. In certain eases the volumes have appeared with considerable irregularity; in others it has occasionally happened that volumes begun in one calendar year have extended into the next year, even when this has not been the general habit of the series. To complicate matters still further, the title-pages in some of these latter volumes bear the later date—a most illogical procedure. In such eases the volume number appears in the accompanying columns opposite both years. In a short summary of this kind it is impossible to give full details in each ease, but the foregoing remarks will serve to explain several apparent anomalies.

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<sup>\*</sup> First series knewn as Bulletin de l'harmacie.

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\* See footnote, p. xx11

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\* Remainder of vol. 55 appeared in 1920.



## A TEXT-BOOK OF INORGANIC CHEMISTRY.

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VOL. IV.

## ALUMINIUM AND ITS CONGENERS, INCLUDING THE RARE EARTH METALS.

#### CHAPTER 1

#### INTRODUCTORY.

RETAINING Mendelcell's original numbering for the series, Group III of the Periodic Table may be represented as in the subjoined table. Buron is the so called "typical element. Buron and aluminium are members of the short periods. The remaining elements belong to the long periods, and

may be divided into the odd subgroup, or Subgroup III. B, viz., Ga, In, and Tl, and the even subgroup, or Subgroup III. A, viz., Se, Y, L., and Ae Aluminium exhibits a marked resemblance to the members of the odd subgroup, with which it is usually classed. Actimum will not be discussed in the present chapter.

	-	I
i	1	ı
1	i	1
2	В	
1 2 3 4	· A1	
4	Se	
5	· Ga	٠
- 6	Y	
7	- La	
8	Las	:
0		i
10	?	
11	1 TI	1
12	i Ac	i
L		i

As a non-metal, boron possesses little in common with the other elements of this group, but exhibits a formal valency resemblance, since, like the others, it can function as a tervalent element. The sesqui oxide of boron, however, appears to resemble the sesqui oxides of aluminum, gallinin, and indimir in being amphoteric. Boron also resembles the metals of Subgroup 111-B in forming alkyl derivatives. Apart from the valency difference, there is considerable chemical smalarity between boron and carbon, and even more between boron and silicon.

The metals of the even, like those of the old, subgroup are tervalent, and hence corresponding compounds of all these elements have similar formulæ. The similarity, however, seldom extends to the water of crystallisation in the case of salts, and, except in the cases of the ethylsulphates and

 $<sup>^{\</sup>bullet}$  1 Organometallic compounds of aluminium, indium (Thiel and Koelsch, Zeitsch amorg. Chem., 1910, 66, 288), and thalloun are known, but no attempts have been made to produce gallium compounds of this type

acetylacetonates of scandium and indium,1 compounds of the elements of the even subgroup are not known to be isomorphous with those of the elements of the odd subgroup. Thus, in the even subgroup double sulphates with the alkali sulphates are known, but they are not of the alum type,  $M_2SO_4M_2^m(SO_4)_4$  24H<sub>2</sub>O<sub>4</sub> given by aluminum, gallium, indium, and possibly by thalloun

The hydroxides in Subgroup III. A are much stronger bases than those in Subgroup III. B. With the exception of thallic hydroxide, the latter are amphotetic. In Subgroup III. A the anhydrous chlorides, broundes, and rodides are only slightly volatile at a red heat, in Subgroup III B they are readily volatile. Organometallie compounds are not known in Subgroup III A, but they are in Subgroup III B. In fact, in these and other characteristics, the old and even subgroups of Group III and Standard to the corresponding subgroups of Group II

In various respects scandium differs considerably from yttrium and lanthanum? The latter, however, exhibit a very close chemical resemblance Moreover, they are very similar in chemical behaviour to the rare earth elements, with which they are generally classed. The general chemical characteristics of yttrium and lanthanium are therefore discussed later when dealing with the rare earth elements 3

A number of physical properties of the metals of Subgroup III B are given in the accompanying table, the data for boron being also given for purposes of comparison. It will be noticed that the order of atomic weights is not the order of the melting points, and that aluminimor is anomalous in that it is paramagnetic.

	В	<b>A</b> 1	Ga	l In	11
Atomic weight Density Atomic volume Melting-point Boiling-point Latent heat of fusion (cals per gram) Specific heat Atomic heat Coefficient of expansion 10 Atomic refraction 4 Magnetic susceptibility per gram > 10 Molecule, in vapoin ,, in Hig solution ,, in Sir solution	11 0 2 31 4 72 c 2200 , 0 3066 (0 100 3 37 t 5 4 0 65 3	27 1 2 70 10 03 658 ( 1800) 70 80 0 218 (0 100) 5 91 2 14 9 5 1 0 6 ( Al <sub>2</sub>	69 9 5 96 11 73 30 2° 19 1 0 070 (12 23) 7 14 8 0 23	111 8 7 28 15 77 165 , 0 0 570 (0 100 6 51 1 5 5 17 4 0 1 2 1 10 1 10	201 0 11 85 17 21 302 1300 1500° 7 2 0 03 36 2) 100 ) 6 05 3 14 21 6 0 0 22 11 11 11

The line spectra of aluminum, gallium, indum, and thallium resemble one another in containing doublet series of the principal, diffuse, and sharp

Jacger Proc K Akad Wetensch Amsterdam, 1914, 16, 1095, Rec trace chem., 1914,
 33, 342 See Chapter IA.
 4 Ghabstone and Dale's formula, 11, line
 5 See Owen, Proc K Akad Wetensch Amsterdam, 1911, 14, 637, Ann. Physik,
 1912, [iv.], 37, 057.

types 1 The sharp series are the best known of them. Each may be represented by a lineks equation —

$$S(m) = Sx - 109675 (m + \mu + a_1'm)^2$$
.

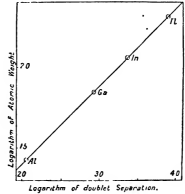
The values of the constants are as follows = 2

				, ,			
Me tal	Atonne Wright W	S <sub>1</sub> z	μ	α.	.	$W^2/\nu$ .	
					: - I		
Al Ga In	27 1 69 9	18161 16 16415 16	1 250525 1 292185	- 0 063319 - 0 077314	112 15 826 10	6 54 5 91	
i ii	1118 2010	11151 76 11170 23	1 287179 1 261328	= 0*069510 = 0 070417	2212 38 7792 39	5 96 5 31	

Here r denotes the doublet separation, or, in other words, the number to add to  $S_1 z$  to obtain  $S_2 z$ . It is obvious that the values of the doublet separations r are roughly proportional to the squares of the atomic weights W.

They are, however, much more nearly in accordance with Runge and Pricht's law that the logar ithms of  $\boldsymbol{\nu}$  are proportional to the logarithms of W . The relation ship is shown graphically in fig. 1 It is of interest to note that the line spectra of scandium, yttimin, and lauthanum also contain doublet series of the sharp and diffuse types 1

Mummum and gallmm com pounds impart no colour to the Buusen flame, the spark spectrum of gallium is, however, very sensitive. Indom and its compounds colour the flame dark blue, thallium and its compounds a bright green, the flames show well defined spectra-



Relationship between spectra and atomic weights of Al, Ga, In, Tl 1'16. 1

The electrode potentials of aluminum, gallium, indium, and thallium are not known with much accuracy, but aluminum is the most electropositive of these elements. The metals alummum, gallinin, and indium pass into solution in dilute numeral acids as tervalent ions Al., Ga., and In., but thallium yields the univalent ion TF.

The chlorides of the metals of Subgroup III. B are as follows -

For a short account of ser●s lines in spectra, see Vol. I. of this series
 On the cries lines in the spectra of Al, Ca., In, Tl, see Hicks Phat. Trans., 1912, A,
 33., 1913, A., 213, 323., Paschen and Meissner, Ann. Physik, 1914, [iv.], 43, 1223
 See Watts, Phat. Mag., 1906, [vi.], 8, 279

The ability of the metals to form chlorides increases with increase of atomic weight. It is interesting to observe that whereas dry hydrogen chloride reacts with heated aluminum and gallium to give AlCl<sub>3</sub> and GaCl<sub>3</sub>, it yields InCl<sub>2</sub> and TlCl with indium and thallium respectively. The trichlorides are hygroscopic solids of low melting-point and volatilise at comparatively low temperatures. They are accordingly intermediate between the less volatile dichlorides of the metals of Subgroup II B and the more volatile tetrachlorides of the elements of Subgroup IV B. The stabilities of the trichlorides diminish in the order Al, Ga, In, Tl. Aluminium chloride is stable at a bright red heat, gallium trichloride begins to dissociate at c. 1000°, and indium trichloride at a rather lower temperature, while thallium trichloride dissociates at temperatures below 100°. The bronides of the metals of Subgroup III, B closely resemble the chlorides.

Each of the metals under discussion forms a sesqui oxide. The stabilities of the sesqui-oxides diminish in the order of increasing atomic weights of the metals. Thus, hydrogen at a bright red heat has no action on alumina, but reduces gallium sesqui oxide to the metal, the reduction of indium sesqui-oxide is easier to accomplish, and the reduction of thallic oxide still easier Moreover, the last named sesqui oxide loses oxygen when heated, passing into thallous oxide.

The hydroxides, M(OH)<sub>p</sub> are feeble bases, practically insoluble in water lt is difficult to say definitely what is the order of their relative strengths, but gallium hydroxide is undoubtedly a weaker base than the hydroxides of alumnium and indium. With the exception of the thallic compound, the hydroxides are also teeble acids. Here, again, gallium appears to be anomalous, in that its hydroxide is a stronger acid than the hydroxides of alumnium and indium.

The soluble salts of aluminum, gallium, indium, and thallium,<sup>2</sup> being salts derived from weak bases, are hydrolysed appreciably in aqueous solution at the ordinary temperature, and the degree of hydrolysis increases with rise of temperature. Normal salts of very weak acids, eg carbonic acid, cannot be prepared

The sesqui-sulphides,  $M_2S_3$  are of interest. Aluminium sulphide is completely hydrolysed by water. Gallium sulphide can be precipitated from gallium salts in acetic acid or animounical solution, but only in the presence of another sulphide. Indium sulphide can be precipitated even in the presence of immerial acids provided that the concentration of acid is very small. Finally, thallie sulphide cannot be obtained by double decomposition in aqueous solution since it is readily reduced to thallous sulphide.

The double sulphates of the type M2SO<sub>4</sub> M2SO<sub>4</sub> M2SO<sub>4</sub> 24H<sub>2</sub>O, where M<sup>1</sup> - Na, K, Rb, Cs, NH<sub>2</sub> or Tl, and M<sup>21</sup> - Al, Ga, Iu, or Tl, are known as alums, and form a group of isomorphous salts. Lathium alums are not known with certainty, although from the nature of the known sodium alums it is reasonable to suppose that lithium alums might exist at temperatures below the ordinary, say O.C. The sodium and potassium indium alums are not known. The only thalke alum known is the ammonium salt, and that has only been obtained as mixed crystals in association with the aluminium salt. Thus, as the atomic weights increase, the metals of the old subgroup form alums with increasing difficulty.

<sup>&</sup>lt;sup>1</sup> Lecoq de Borsbandian , see "Gallium" in Wurtz, Dationnaire de chimu, Supplement, p. 859.

<sup>2</sup> Thallic salts are understood here.

<sup>3</sup> See p. 82.

In addition to the alums, double sulphates of the type M So<sub>4</sub> M<sub>2</sub> (SO<sub>4</sub>)<sub>3</sub>, 811,0 are known. The following have been prepared -

```
In (804), (NII4) 804 SH O
In (804) Na 804 8H O
In (804), K(804 SH O
\begin{array}{l} Al_2(8O_4), \ (NH_4), SO_4, 8H_2O \\ Al_2(8O_4), \ K_2SO_4, 8H_2O \end{array}
                                                                                                                                                                                                                                                                                                      TI (SO<sub>4</sub>) (NH<sub>1</sub>\2SO<sub>4</sub> 8H<sub>2</sub>O<sub>4</sub> TI (SO<sub>4</sub>) | K (SO<sub>4</sub> 8H<sub>2</sub>O<sub>4</sub> TI, SO<sub>4</sub>\2 Rb<sub>2</sub>SO<sub>4</sub> 8H<sub>2</sub>O<sub>4</sub>
```

With these salts the stability increases with increase in atomic weight of the tervalent metal present. The aluminium salts, for instance, are not stable in contact with aqueous solutions. In the case of the indium ammonium salt it is known that the reversible change

$$In_2(SO_1)_3 (NII_1)_2SO_1 24II_3O_2 - In_3(SO_1)_3 (NII_1)_2SO_1 8II_2O + 16II_3O_2$$

occurs, the transition point being 36 C. It is therefore very probable that the sodium and potassium indium alums could be prepared at low temperatures. The octahydrates are of interest owing to the fact that several sulphates of rare earth elements form double salts of the same type

The alums afford instances of isomorphism between compounds of aluminium, gallium, indium, and thallium. Few other cases are known. The two following isomorphous series have, however, been examined in detail. detail

The diteti igonal bipyramidal series, R-MN<sub>6</sub> 211.0.

2. The thombse bipyramidal series, R.M.V. 11.0

The isomorphism between the double chlorides and broundes of tervalent indium and thallium is valuable evidence in favour or classifying these two metals together in the same natural lamily

Little is known of the compounds or bivalent gallium. The halogen compounds, however, like the halides of univalent and lovadent indium, appear to possess few of the properties of salts. Univalent thalloun, on the other hand, forms a well-defined series of stable salts, while the compounds of tervalent thallour are ill defined and unstable. In this and in other ways thallum differs from the other members of Subgroup III, B, the pecularities of this element, however, will be discussed later?

The corresponding double seterate is also known
 Wallace, Leitsche Krust Min., 1911, 49, 417
 See Chapter VIII.

#### CHAPTER II.

#### BORON.

Symbol, B. Atomic weight, 11.0 (O = 16).

Occurrence -Boron is widely distributed in nature, but it never occurs in the free state. It is present in minute amounts in sea water 1 and mineral waters,2 and is widely diffused in the vegetable kingdom,3 the ashes of nearly all fruits contain small quantities of boron, which is accordingly found in wines. Boron has been detected in the organs and tissues of various animals. It is also a constituent of certain numerals, of which tour matine, danburite, and armite may be mentioned

Small trichme crystals of boric acid are occasionally found in nature, being known as sassolite. Boric acid is also present in the jets of steam that issue from the earth in certain volcame regions, and which are called suplion in Tuscany Salts of horic acid occur in various parts of the globe, notably in the Great Basin region of the western United States, in Chili and Peru, at Stassfurt in Germany, in Asia Minor, and in Tibet. The most important deposits are in the United States, where, after borar had been found in the water of Clear Lake, California, vast surface deposits of borax were found in the desert marshes of California and Nevada. In 1890, large deposits of colemanite were discovered, embedded in old tertiary sediments

Mmeral barax, Na<sub>2</sub>B<sub>3</sub>O<sub>7</sub> 10H<sub>3</sub>O, forms monochine prisms of density 1.7 and hardness 2 to 25". It was originally obtained from a salt lake in Tibet and sent to Europe under the name of tinkal

Colemanite (or borate spar) is a hydrated calcium borate of the formula  $\text{Ca}_2\text{B}_6\text{O}_{13}5\text{H}_2\text{O}$  . In appearance it resembles calespar, hardness, 3.5 to 1.5, density, 2.4. Two other varieties of calcium borate are also tound. *priceite* (or bechilite) and pandermite Priceite occurs in the United States in County Oregon, and is a fine, white, soft, chalky mineral Pandermite occurs in an enormous bed of gypsum covered with several feet of clay, in Asia Minor, just south of the Sea of Marmora. It is exported from the port of Panderma. In appearance, pandermite closely resembles a fine grained white marble

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 12, 318., Campt. rend., 1881, 93, 224., 1882, 94, 1352
 2 Goodh and Wintheld, Bull. U.S. Geol. Survey, 1888, No. 47., Fonzes Diacon and Fabre,

Compl. rend., 1914, 158, 1511

<sup>3</sup> Jay, Compl. rend., 1895, 121, 896. For hammer, Phil. Trans., 1865, 155, 203, Crampton, Bra., 1889, 21072, Baumert, Bra., 1888, 21, 3290, Bechn. Bull. Soc. chim, 1890, [10], 3, 122; Hotter, Chem. Soc. Abst., 1890, p. 1398, Passerim, thid., 1893, u. p. 225; von Lippmann, Chem. Zeit., 1902, 26, 405, Avzarello, Gazzetta, 190a, 36, n. 575, Dugast, Compl. cend., 1910, 150, 838; Bertainchand and Dugast, Ann. Chem. Anal., 1910, 1519 1910, 15, 179.

<sup>4</sup> Jay, loc. cit.; Bertrand and Agulhon, Compt rend , 1912, 155, 218, 1913, 156, 732.

Ulcarte (beronatrocalcite, to a, or cottonballs) is a sodium calcium borate, Na B<sub>3</sub>O<sub>2</sub> Ca B<sub>3</sub>O<sub>4</sub> 16H<sub>3</sub>O<sub>5</sub> which occurs in considerable quantities in Peri and Club. Hardness, L., density, 1.65. It is found as yellowish white nodules, often incrusted with sodium sulphate and chlorole

Boracite (or  $stassimitite), <math display="inline">2{\rm Mg}^2$  B, O  $_{\rm Le}{\rm MgCI}$  , is a double borate and chloride of magnesium. It crystallises in the fetrahedral class of the regular system, hardness, 7, density, 30. Boracite is found at Stassfurt embedded in the kainite deposits 1

History .-- Boray has been known in commerce for many centuries. The name occurs in the writings of Geber and other abhemists, but it is difficult to say whether they refer to the boray of to day. In 1702 Homberg? prepared boric acid from boacs, and called it sal solutions. Temers, in 1728, showed that sal-sedativiiii could be obtained by adding any immeral acid to boray, and Baron, in 1747,1 discovered that boray was composed of sill solutivum and solut. With the establishment of Lavoisier's system of nomenclature, the name sal solativum gave place to bordere will, and the acid was looked upon as the oxide of an unknown element. This element, horon, was isolated in an impure state in 1808 by Gay Lussae and Thenard, and by H. Davy

Preparation. The tollowing methods have been proposed for the isdation of boom

- (r) Reduction of boron sesqui oxide with potassium, sodium, magnesium, According to Wohler and Deville, 10 parts of boron sesqui or aluminum oxide are mixed with 6 parts of sodium, covered with 5 parts of sodium chloride, and heated to redness. The magnesium process is discussed later
  - (ii) Electrolysis of fused boron sesqui oxide for horax 10
  - (m) Reduction of Loron trilluoride by means of heated potassium  $^{\rm H}$
- (iv.) Reduction of potassium bondhoride by means of heated potassium  $^{19}$ or magnesium 13
  - (v) Reduction of loans by heated magnesmin 11 or red phosphorus 15
- For brither information concruming the eccurionic of boson, see Merrill, The Non-Metallic Memorals, 2nd of (Wiley & Sons, 1910), p. 231 of Laumay, Trade de Metallicanic, 3 vols. Paris, 1913, vol a p. 6.4, Impoint J. Frag. Ind. Chem., 1910, 2, 500; Warnswright From Marche by Gol. Min. Soc., 1909, 31, 601. Reachett Com. Zoule, 1910, p. 197; Bocke, Cod. Min., 1910, p. 541; Keyes Fing Min. J., 1940, 88, 826.
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   Barun Mon. Ind. Soc., 17, 1750, r. 1295, 147.
   Gry Los wand Themaid C. B. Acc. at n., 1808, 30, 363. Ann. Chem. Phys., 1809, 69, 204., Restar her phus. archimage. vol. 1, p. 276.
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  Rawson, Chem. Xeit., 1888, 58, 284.

- \*\*Cittle Harimana, Sett at Science 1, 1944, 2, 34.

  \*\*Barrahus, Poqo Inno (20, 124, 2, 113)

  \*\*Garther, Jahrester 1, 1865, p. 126

  \*\*Gartermana, Rev., 1889, 22, 195, Marsch, Chem. Zentr., 1889, u. p. 905, Ray,
- vide supra <sup>15</sup> Diagendorff, Chem. Zentr., 1861, p. 865.

- (vi.) Reduction of boron trichloride by means of hydrogen 1
- (vii.) Reduction of the oxide, brounde, or intride of boron by heating with metallic calcium; 2
- (vin) Dissociation of magnesium boude, "boron suboxide," and boron intride in vacuo at high temperatures (1500° and above) <sup>3</sup>

Pure boron was first obtained in 1909, by Weintranh <sup>1</sup>. Previous to this, Moissan claimed to have prepared boron containing not more than 2 per cent of impurities <sup>1</sup>. According to Moissan, the early processes described by Guy-Lussac and Themard, Berzelins, and Wohler and Deville, lead to products which, after puriheation by heating with water and hydrochloric acid, only contain 40 to 70 per cent of boron.

According to the method adopted in its preparation, pure boron may be obtained either in small grains having a sub-metallic appearance or in the massive form, the lumps exhibiting a conchoidal fracture. It is very probable, but not certain, that in each of these forms boron is crystalline. The so-called crystalline varieties of boron described by the earlier chemists are, however, bondes of aluminum.

Moissan, who considered that he had prepared nearly pure boron, described it as an amorphous, brown powder, and most of the carber investigators described it in similar terms. The researches of Weintraub and of Ray have shown, however, that boron obtained by Moissan's method contains several per cent of ovegen and that under certain conditions the oxygen content may reach as much as 16 per cent. In fact, the reduction of excess of either boron sesqui-oxide or borax with a sintable metallic reducing agent always leads to the production of an amorphous, brown powder containing oxygen "Weintraub refers to such a product as boron subaxide, it is most probably a solid solution of an oxide of boron, possibly  $B_4 O_{\rm go}$  in boron? This so called amorphous boron is rapidly converted into boric acid when gently warned with 10 per cent intricated. Under such treatment pure boron is hardly attacked. The small proportion of pure boron usually present in the amorphous product can therefore be readily isolated.

The practical methods for obtaining pure, or nearly pure boron, are as follows —

1 Reduction of Boson Trifluoride -- The fluoride, generated in a lead or platinum apparatus to exclude silicon fluoride, and free from hydrogen fluoride, is led over sodium heated in a hard glass tube to the lowest temperature at which reduction can be effected. The product is extracted

 $<sup>^1</sup>$  Weintraub, vide infra . Pring and Fielding, vide infra . Bessen and Fourinei, vide infra.

<sup>&</sup>lt;sup>2</sup> Moissan, Ann Chim Phys., 1899 [vn.], 18, 289, cf., however, Stock and Holle, Bur., 1908, 41, 2095

Wentraub Trans, Amer. Electrochem. Soc., 1909, 16, 165., J. Ind. Fau. Chem., 1911,
 299., 1913, 5, 106., U.S.A., Pat., 997,879; 997,880, 997,881, 997,882, 1,019,394,
 1,019,569; Eng. Pat., 21,667, 23,334, 25,978 (1906), 1197 (1907), 25,033 (1910), 7103 (1911).

<sup>&</sup>lt;sup>4</sup> Mossan, Compt. rend., 1892, 114, 319, 392. Ann. Chim Phys., 1895, [vn.], 6, 296, see also Binet du Jassonners, thal., 1909, [vm.], 17, 145

<sup>&</sup>lt;sup>5</sup> See p. 97.

<sup>6</sup> The product obtained by the electrolysis of fused borax also appears to be amorphous and to contain a considerable quantity of oxygen.

<sup>7</sup> Ray, 11de supra.

with water and hydrochloric acid in a platinum dish, when boron is left as a brown powder 1

A more convenient process consists in substituting potassium borothioride for boron trifluoride. Excess of the potassium salt is mixed with sodium in a crucible lined with magnesia, and heated to redness in a rapid stream of dry hydrogen. The boron is isolated as before

2 Reduction of Boson Sesque orde with Magnesium and further Reduction of the "Suboxide" obtained - Ently experimenters used for this purpose 2 parts of oxide to 1 of metal, the reaction being considered to be as follows ---

$$2B_2O_3 + 3Mg = Mg_3(BO_3)_2 + 2B$$

Moissan increased the amount of oxide to 3 parts, thereby keeping the "amorphous boron" produced suspended in a very fluid mass of boron sesqui-oxide and away from the an-His method is as follows - 2

Boron sesqui oxide (210 grams), free from sodium, calcium, and silieou compounds and recently fised and powdered, is mixed in an earthenware crueible with magnesium turnings (70 grams) free from non and silicon. The criterible is covered and placed in a furnace already at a bright red heat Within five minutes, a violent reaction takes place. The mass is heated for another ten minutes, then cooled and removed from the erneible. Externally the reaction product is black internally it has a maroon colom, traversed in all directions by white crystals of magnesium borate. The black outer layers are removed and rejected. The residue is powdered and boiled with a large quantity of dilute hydrochloric acid. It is then heated with pine hydrochloric acid at the boiling point of the latter for two hours, a procedure that is repeated six times. The solid residue is washed with water, licated with boiling 10 per cent, alcoholic potash, and again washed with water. Finally it is heated in a platinum dish for four hours with 50 per cent. hydrofluoric acid, washed with water, and dired, first on a porous plate and then in a vacuum over phosphoric anhydride. The product contains 88 to 90 per cent. boron and 2 to 1 per cent magnesium (Weintranh), the preceding treatment having served to free it almost completely from boron sesqui-oxide, magnesium borate, boron intride, and silica. To eliminate the remaining magnesium, the boron is mixed intimately with hity times its weight of boron sesqui oxide, introduced between two layers of the oxide in a large earthen erneible and heated to bright redness. The product, when cold, is treated by the process already described. In this manner it is possible to obtain a brown, amorphous powder containing only 0.4 per cent of magnesium. It is advantageous, during each heating in the furnace, to enclose the criterible in a larger one brasqued with a finely powdered mixture of rutile and carbon. An alternative method of chimmating introgen is to carry out the preparation in an iron vessel lined with magnesia, a rapid stream of hydrogen being passed through it 1

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<sup>1</sup> Prof. H. Breicton Baker, private communication. Rawson, vole supra.

Monsan, cute supra
 This magnesium is present neither as borate nor boride, but is present in solid solution either as magnesium oxide or possibly as a borite (Ray)
 Considerable purification may quickly be effected if, after the products of the interaction of boron sesqui-oxide and magnesium have been treated with hydrochloric acid, the residue is submitted to fractional decaptation and the lighter fractions discarded (Brief du Jassonners, vide supra, Weintiaub, vide supra)

In the course of his investigations of the magnesium process, Weintraub 1 found that if a higher temperature than that used by Moissan is employed, there is no difficulty in obtaining a product free from magnesium. To obtain the higher temperature, Moissan's procedure may be used, with charges of at least one kilogram of material, or the reaction may be effected in a graphite crueible heated to 1700° in a vacuum electric furnace. The high temperature

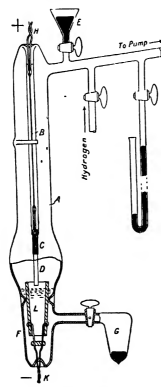


Fig. 2. - Mercury are furnace for melting boron.

product, however, contains not 4 to 5 per cent but 14 to 16 per cent of oxygen, and re-insion with a large excess of boron sesqui oxide is without action upon it. If the ratio oxide, metal is diminished below 2. I, the product obtained, after the usual chemical treatment, consists mainly of magnesium bonde.

Magnesium boride, "boron suboxide," and boron intride all dissociate when heated in vacuo to sufficiently high temperatures With the boulde, the change is perceptible at 1200°, and is rapid at 1500°. The subovide and mitride, however, dissociate with more difficulty, particularly the latter Owing to this dissociation, it is comparatively easy to obtain pure boron from the so called "amorphous boron" or "boron suboxide," and at the same time to melt the boron (1) One method consists in placing the powder in a water cooled copper cup which forms one of two copper electrodes, and passing an arc between them in vacuo or in hydrogen at a few millimetres pressure Several pounds of boron may be prepared and melted at one operation in this manner, starting with boron suboxide (Weintraub). (ii ) Another method consists in using a "mercury are furnace" In this case, Moissan's boron or boron suboxide should be compressed into rods and purified as far as possible by heating

to 1200° to 1500° in vacuo. A boron rod D, mounted in a carbon holder C to which a lead B is attached, is placed within the cylindrical glass apparatus A (fig. 2). The apparatus may be filled with hydrogen or evacuated by means of suitable connections. The apparatus being evacuated, mercury is run in through the funnel E until the end of D is just covered, Hydrogen is then admitted up to 2.5 to 15 cms, pressure, the electrodes H and K connected to a suitable source of current, and mercury run into C until the end of D is clear of the mercury in the lower part F of the

<sup>1</sup> Weintraub, vide supra

apparatus. An aic is thus formed between the boion and the mercury in , and, since at the surface of the latter great heat is developed, the end of D melts and falls off. As the rod D thus shortens, more mercury is run in from time to time from E. The bottomless alundum pot L prevents the are from straying to the glass. In this manner, extremely pure boron may be obtained (Weintraub) (iii) A third method, somewhat analogous to the preceding method (n), consists in focussing the cathode rays on the powdered material, an operation that must obviously be carried out in a nearly perfect vacuum 1

The preceding methods for obtaining pure boron necessitate the employment of special forms of apparatus. A fourth method is very simple to carry out, but does not give the product in the massive form (iv) The "suboxide" is mixed with excess of magnesium or sodium and heated to (iv) The bright redness in a stream of hydrogen, in the former case magnesium bonde is formed and subsequently decomposed at the high temperature employed. The product is extracted successively with water, hydrochloric acid, and warm 40 per cent intric acid '

3 Reduction of Boron Trichloride with Hydrogen -This method, also due to Weintraub, is based upon the reversible reaction

$$2BCl_3 + 3H_2$$
.  $2B + 6HCl$ ,

which has been studied by Pring and Fielding, and also by Besson and Fournier,3 A high temperature is necessary for the production of boron by this method, and is obtained by the use of an electric are. The preparation is effected by running one or more alternating current ares between water or an cooled copper electrodes in a mixture of boron trichloride vapour and a large excess of bydrogen, a glass or copper contaming vessel may be used. The gases are inixed by passing dry hydrogen over the surface of boron trichloride contained in a vessel cooled by a freezing mixture, or by allowing the chloride to drop into a vessel (kept a little above the room temperature) through which hydrogen is passed. The boron is partly thrown off as a line powder and partly settles on the electrodes, where it grows into small rods. These eventually melt down to beads and fall off, and the process repeats itself The boron powder, after washing with water, contains 99 to 99 5 per cent of boron, the fused lumps are pure boron. By any of the processes aheady mentioned the boron powder may be purified and melted

Properties, -- Boron is a black solid of density 2:34 and compressibility 0 3 10 6 per atmosphere at 20 4 In hardness it is interior only to diamond. Its fracture is conchoidal, it shows no signs of microcrystalline structure It is inferior to diamond in its toughness or strength. Under atmospheric pressure boron melts at about 2200 5. It passes into vapour at temperatures considerably below the melting-point, the vapour tension becoming noticeable at 1600. The electrical conductivity of a piece of massive boron at ordinary temperatures is extremely small, but between room temperature and that corresponding to a dull red heat it increases to \_\_\_\_\_\_

<sup>1</sup> Tiede and Bunbauer, Zeitsch annry Chem., 1914, 87, 129, Tiede, Ber , 1913, 46,

Ray, Trans. Chem. Soc., e1914, 105, 2162
 Pring and Fielding, Trans. Chem. Soc., 1910, 95, 1497, Besson and Fournit, Compt.
 Pend., 1910, 150, 872, Weintraub, Ion. ext.
 Richards, J. Amer. Chem. Soc., 1915, 37, 1646
 Tiede and Biribrauer, Zeitsch. anorg. Chem., 1914, 87, 129, Weintraub gives 2800°.

about two million times its initial value. For instance, a piece of boron which had a resistance of 5.62 × 106 ohms at 27° had a resistance of only 4.60 101 ohms at 180° (1 At a dull red heat the resistance had fallen to 5 ohms. Over a short temperature interval the resistance of boron is an exponential function of the temperature. At 23° the specific resistance is about 1.7° 10° ohms per cm. cube; at 0° the value is about 2° 10°. The specific resistance of boion is enormously decreased at ordinary temperatures by the introduction into it of small amounts of other elements, and at the same time the very large negative temperature-coefficient is diminished Thus, a few tenths per cent of carbon introduced into boion reduces the resistance to about one-twelfth its value at ordinary temperatures 1

The atomic refraction of boron in saturated compounds for the D line is 5.3 or 3.2, and for the H<sub>a</sub> line 5.4 or 3.1, according as the Gladstone and Dale or the Lorenz and Lorentz formula is employed 2

The spark and are spectra of boron consist simply of three lines in the ultraviolet, 3451 50, 2497 83\*, and 2496 89\*, the pair asterisked being more persistent than the other line. Two band spectra, however, are associated with boron. The green flames due to boric acid, alkyl borates, and boron fluoride all have practically identical band spectra, the bands are headless, and the most prominent are in the green region (see fig. 3 on p 44). They have been observed and measured by numerous scientists <sup>4</sup> The same bands are observed in the are spectra of boron and boron sesquioxide, and in the spark spectrum of a solution of boric acid in hydrochloric acid, and they are attributed to oxide of boron, they do not occur in the spark spectrum of boron itself. Another series of bands, which have definite heads and degrade towards the red, is found in the spectrum of boron trichloride or methyl borate in the afterglow of active introgen (p. 24), these bands also occur to some extent in the arc spectrum of boron and its oxide, and are attributed to boron mittide but they do not occur in the  ${\bf spark}$  spectrum of boron m introgen  $^6$ 

Pure boron may be strongly heated in air without undergoing any perceptible oxidation. It is oxidised to bone acid when heated with concentrated intric acid, but the rate of oxidation is very slow. It does not combine with either copper (Weintraub) or magnesium (Ray) at a red heat. Its other chemical properties are unknown

Moissan's "amorphous boron" is a maroon coloured powder of specific gravity 2.45. Its specific heat increases rapidly with use of temperature, the

<sup>1</sup> Weintraub, ride supra

Wentraub, side supra
 Ghus, Gazzetta, 1893, 23, 1-452, n. 8.
 Crookes, Proc. Roy. Soc., 1912, A, 86, 36, de Gramont, Compt. rend., 1908, 146, 1260; cf. Hartley, Trans. Chem. Soc., 1883, 43, 390; Proc. Roy. Soc., 1883, 35, 301, Rowland, Phil. May., 1893, [v.], 36, 49; Knyser and Runge, Wod. Annalen, 1893, 48, 126; Eder and Valenta, Denkschr. K. Akad. Wes. Wea, 1893, 60, 307; Evner and Haschek, Stang-ber. K. Akad. Was. Wien, 1897, 106, 11a, 494, Kaysen, Handbuch der. Societoschene (Lenna, 1910), ad. p. 134.

Haschek, Steing-ber. A. Akad. II is. Wien, 1897, 100, 11a, 494, Kaysei, Hanabuch der Spectroskopie (Leipiig, 1910), vol. v. p. 134.

Leeoog de Boisbaudium, Compt. rend., 1873, 76, 833. Spectres Lumineux (Paiis, 1874), Salet, Ann. Chim. Phys., 1873, [iv.], 28, 5, Traité de spectroscopie (Paiis, 1888); Dieulafat, Ann. Chim. Phys., 1877, [v.], 12, 318; Chamleian, Silzangsber. K. Akad. Wiss.

Disulafat, Ann. Chim. Phys., 1817, [v], 12, 310; Chambelan, Steadyson.

Wiem, 1880, 82, II, 425.

Lecoq de Boisbaudran, loc cit., Hagenbach and Koren, Atlas der Emission-yektra
(Jena, 1905); Hagenbach, Wuilner Festschrift (Leipzig, 1905), p. 128, Kuhne, Zeitsch.
Wiss. Photochem, 1906, 4, 173, Anerbach, ibid, 1909, 7, 30, 41; Eder and Valenta,
Denkschr. K. Akud. Wiss. Wien, 1893, 60, 467, Jevons, vode infra.

Bevons, Proc. Roy. Soc., 1915, A, 91, 120; cf. Kuhne, loc cit.

mean value being 0 3066 between 0° and 100°, 0 3407 between 0° and 192°, and 0.3573 between 0° and 235°.1

The chemical properties of "amorphous boron" have been described in detail by Moissan," and are given in the following paragraphs. It must, however, be remembered that these properties refer to a substance that has been shown by Weintlaub to contain 1 to 5 per cent. of oxygen It will be noticed that several of the properties mentioned are not the properties of pure boron.

Boron unites directly with fluorine at the ordinary temperature, with chlorine at 410°, and with bromme at 700°, but is without action on iodine. With the halogen acids it reacts with greater difficulty, hydrogen fluoride attacks it at a dull red heat, hydrogen chloride at a bright red heat, the products being hydrogen and boron fluoride or chloride. Hydrogen nodide has no action on boron. When heated in oxygen, boron ignites with the production of heat and light, and it burns in air at 700'3. It umtes readily with sulphur at 600°, producing the sesqui-sulphide, and combines with selenium at a higher temperature, it does not combine directly with tellurium The direct combination of boron and nitrogen occurs very slowly at 900°, but rapidly at 1250°. Neither phosphorus nor arsenic vapour at 750° will combine with boron, and antimony may be fused with it without the production of any chemical change. When heated with carbon in the electric are in an atmosphere of hydrogen, crystalline boron carbide is produced.

Boron does not unite directly with the alkali metals - On the other hand, it unites with magnesium at a dull red heat, and with iron and aluminium at a higher temperature—It also mates with silver and platinum

Boron acts as a powerful reducing agent. It decomposes steam at a red heat, reduces iodic acid on warming, with the liberation of iodine, and at a temperature just below a dull red heat it reduces sulphur dioxide. At a dull red heat the oxides of arsenic and carbon dioxide are reduced by boron, at 800' phosphoric anhydride is reduced, and at 1200° the reduction of carbon monoxide and silica can be effected. At a bright red heat boron burns in either introus or intric oxide, producing boron oxide and intride. Concentrated sulphune and intrie acids are readily reduced by boron, which becomes oxidised to boric acid

A large number of oxides and salts are reduced by boron. The oxides of copper, tin, lead, antimony, and bismuth are reduced when gently warmed with boron, the mixtures becoming meandescent. At a red heat the oxides of iron and cobalt are reduced, but not the alkaline earth oxides. Silver fluoride is violently reduced by boron at ordinary temperatures, the fluorides of zme and lead at a red heat, but not the alkali and alkaline-earth fluorides. Chlorides are less casy to reduce, but mercuric chloride can be reduced to merciny at 700°. When boron is fused with an alkali hydroxide, a violent evolution of hydrogen occurs, and an alkali borate is produced.

Reductions may be effected at the ordinary temperature with certain aqueous solutions. Thus, potassium permanganate is slowly reduced, and

<sup>1</sup> Morssan and Gautier, Ann Chim Phys., 1896, [vii ], 7 568, ct Kopp, F Chem Soc, 1866, 19, 187, F Webet, Phil May, 1875, [iv ], 161, 276, Mixter and Dana, Annalm, 1873, 169, 388, see also Vol I, p 90

2 Morssan, Ann Chim Mays., 1895, [vii ], 6, 296, Compt. rend, 1892, 114, 617; 1893, 117, 423, 1894, 119, 1172

3 The combustion is prevented by carefully drying the elements (Baker, Phil. Trans., 1888, 175, 590)

<sup>1888, 179, 590).</sup> 

precipitates of the respective metals are produced by the action of boron on aqueous solutions of silver nitrate and the chlorides of gold, platinum, and palladium.

Boron will probably receive various practical applications in the near future, owing to its remarkable electrical properties. Its volatility prevents its employment in the manufacture of electric lamp filaments. Boron is of great use in the purification of copper, if 0 03 to 0 1 per cent be added to molten copper before it is east, the metal is deoxidised and purified to a remarkable extent It is cheaper, however, to use "boron suboxide" for the purpose.1 A little boron increases the breaking stress of steel 2

Colloidal Boron. -Colloidal solutions of boron may be prepared from boron obtained by the action of magnesium, sodium, or potassium on boron sesqui-oxide, by extracting the products of the reaction with hydrochloric acid and washing the residue repeatedly with water until the water acquires a decided colour. Two boron hydrosols have been obtained by the magnesium method (1) a relatively unstable sol, and (11.) a comparatively stable sol. The first is dark brown in colour, and may be kept in a closed vessel for not more than three or four weeks. It is immediately coagulated by electrolytes, and rapidly oxidised by dilute intric acid or hydrogen peroxide. It cannot be purified by dialysis. The second resembles the first in appearance, but may be kept for mue weeks and purified by dialysis. Two sols have been obtained by the potassium method, they resemble those just described. Three sols have been obtained by the sodium method, and are deep blue in colour. Two are unstable, but the third is stable and may be purified by dialysis Boron hydrosols have a high electrical resistance, the boron particles are negatively charged

Atomic Weight.—It follows from the analyses and vapour densities of numerous boron compounds that the atomic weight of boron is three times its combining weight

The determinations of the combining weight leave much to be desired Various attempts have been made to measure the ratio Na<sub>2</sub>B<sub>4</sub>O<sub>7</sub> 10H<sub>2</sub>O Na<sub>2</sub>B<sub>4</sub>O<sub>5</sub>, but the results of different experimenters do not agree, and the method, involving as it does the use of a hydrated salt, is a bad one. The best determinations of the combining weight are probably those due to Gantier,4 whose values for the atomic weight of boron, derived by four methods, are as follows -- 5

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∴ B = 11 028
\begin{array}{ll} B_2S_3 & 3BaSO_4 : \\ B_6C & CO_2 \end{array}
                                16.8855 \cdot 100
                                                                 .: B -10 999
                               177\ 258 . 100
\overline{BBr_{s}} + 3\Lambda \overline{gBr}
\overline{BCl_{s}} = 3\Lambda \overline{gCl}
                                 44 512 · 100
                                                                        B . 11 027
                                                                 ٠:.
                                                                .. B = 10.955
                                 27 2845 100
```

The first two ratios are only based upon a few experiments, and from the nature of the analytical operations involved the results cannot be

Weintraub, U.S. Pat., 1,023,601; see also J. Ind. Eng. Chem., 1910, 2, 477
 Guillet, Compl. rend., 1907, 144, 1019. See also Moissin and Charps, Compl. rend., 1895, 120, 130, Hannesen, Zeitsch. anning Chem., 1911, 89, 257, and Vol. IX. of this series.
 Ageno and Bazetti, Atti. R. Accad. Linear, 1910, [v.], 19, 1-381, Gutbier, Kolland.

Zeitsch, 1913, 13 137

4 Gautter, Jun Chim. Phys., 1899, [vii] 1, 18, 352.

5 Clarke, A Readvilution of the Atomic Winghts, 3id ed & "Smithsoman Miscellaneous Collections," vol. 54, No. 3, 1910). The antecedent data have been changed to — Ag=107.880, Cl=35.457, Bi=79.916, S=32.065, C=12.003, Ba=137.37, O=16.000, Na = 22.996.

regarded as possessing much more than corroborative value chemical point of view the other two ratios would be expected to yield more rehable results, but Gautier's method of determining them is open to serious criticism

Mention should be made of Ramsay and Aston's determination of the ratio Na $_2$ B $_4$ O $_7$  2NaCl, effected by distilling anhydrous borax with hydrochloric acid and methyl alcohol  $^1$ . The results obtained lead to the value B 10 951 The other determinations 2 are of little importance

The value at present adopted for the atomic weight of boron is B II'O

## COMPOUNDS OF BORON

From the chemical point of view, boron is decidedly a non-metallic element. Thus, the halides are volatile compounds which are readily and irreversibly hydrolysed by water. The sulphide of boron is similarly hydrolysed by water, several hydrides are volatile. On the other hand, boron sesqui-oxide appears to be amphoteric. From the physical standpoint, boron exhibits a slight resemblance to the metallic elements, the binary systems boron metal, for example, are to some extent similar to the binary systems composed of two metals

From the reactions of boron it is clear that the binary compounds of boron are very stable substances. The heats of formation (H F), given below, fully bear out this conclusion - '

Boron is tervalent in its halogen derivatives, and appears to be usually tervalent in its compounds. However, it seems at times to exhibit a valency of five. Thus, Mossan has prepared boron pentasulphide, and the compounds of the formula Call BCl, (CII), BNII, (Call), BNII, and (C.H. O) B C.H. ONa may perhaps contain pentad boron and be formulated as follows = 5

The compound B(CH), NH, for instance, is fairly stable, it melts at  $51^\circ$  and boils at  $110^\circ$  . The existence of certain additive products of boron

Ramsay and Aston, Trans. Chem. Soc., 1893, 63, 211
 Berzehus, Poop. Annalm., 1826, 8, 1, Lourent, J. prakt. Chem., 1849, 47, 415;
 Hoskyns-Abahall, Trans. Chem. Soc., 1892, 61, 650., Rimbach, Bet., 1893, 26, 164,
 Anintage, Proc. Chem. Soc., 1898, 14, 22
 Troost and Hantefeuille, Ann. Chem. Phys., 1876, [v.], 9, 70., Berthelot, Thermochemic, vol. 2, Statice, Compt. rand., 1891, 112, 862
 Other compounds of this type are known (p. 33)
 Michaelis and Becker, Beg., 1880, 13, 58, Gustavson, Ber., 1870, 3, 426. J. Russ. Chem. Soc., 1888, 20, 621. Ber., 1880, 13, 58.
 Gustavson, Ber., 1870, 3, 426. J. Russ.
 Chem. Soc., 1876, 25, 165. Lorenz, Annalm., 1888, 247, 226. Copaux, Compt., rand., 1898, 127, 779; cf. Frankland, ibid., 1898, 127, 798

trichloride and tribromide also suggests that boron is pentad hypothesis appears necessary to account for the nature of the borohydrates and the lower oxides of boron

The existence of a fairly stable hydride, B2H6, clearly shows that on occasion boron may be at least quadrivalent. According to Stock, boron is quadrivalent in its hydrides, and also in the hypoborates, eg. KOBH<sub>2</sub>, which would accordingly be written

It is of some interest in this connection to note that since boron is occasionally at least quadrivalent, its maximum valency is greater than its group number in the periodic table

Boron Hydrides or Hydroborons -The existence of solid hydrides of boron has been suspected by various chemists. The existence of a volatile hydride was long ago anticipated from analogy with the compounds of other non-metallic elements, but neither Wohler and Deville nor Gustavson succeeded in preparing it. In 1879, F. Jones showed that a volatile boron hydride did exist, and two years later, F Jones and Taylor studied the hydride and assigned to it the formula BH<sub>3</sub><sup>2</sup>. Their work was confirmed in a qualitative way by Sabatiei <sup>3</sup>. Later, Rainsay and Hatfield announced the existence of what were, in their opinion, probably two isomeric hydrides of the formula B<sub>2</sub>H<sub>22</sub> but Ramsay was unable subsequently to duplicate the work. No value can be placed upon the scanty experimental data which served to deduce the preceding formula, since the gaseous boron hydrides were undoubtedly contaminated with silicon hydride

The hydrides prepared by Jones and Taylor, Sabatier, and Ramsay and Hatfield were obtained, mixed with a very large excess of hydrogen, by the action of dilute acids on magnesium bonde, and similar gaseous mixtures can be obtained by the action of acids upon commercial non and manganese bondes 5 From the mixed gases produced from magnesium bonde and hydrochloric acid, Stock and Massenez in 1912 succeeded in isolating two distinct boron hydrides of molecular formula  $B_4H_{10}$  and  $B_6H_{12}$ , and since then Stock and others have obtained several more boron hydrides? Although the compound BII, is not yet known, there are, according to Stock,

Friederici, and Priess, at least ten boron hydrides

Obtained by heating B<sub>4</sub>H<sub>10</sub> Colourless gas (1)  $B_2H_0$ 

(2) B<sub>1</sub>II<sub>10</sub> Obtained from magnesium boride and hydrochloric acid.

(3) A colourless liquid, very unstable, obtained like (1)

(4) B<sub>0</sub>H<sub>12</sub> Obtained like (2).

<sup>&</sup>lt;sup>1</sup> Gattermann, Ber., 1889, 22, 195; Winklei, Ber., 1890, 23, 772, Reinitzer, Silvingsber, K. Akad Wiss. Wien, 1881, 82, 736. Monatsh., 1880, 1, 792, Lorenz, Annalen, 1888, 247, 246, J. Holmann, Chem. Zeit., 1911, 35, 265.

Jones, Trans Chem. Soc., 1879, 35, 41; Jones and Taylor, ibid., 1881, 39, 213.

Sabatiei, Compt. iend., 1891, 112, 8166

Ramsay and Hatheld, Proc. Chem. Soc., 1901, 17, 152.

J. Holfmann, Chem. Zeit., 1911, 35, 265.

Stock and Mussenez, Ber., 1912, 45, 3539

Stock and Friederici, Ber., 1913, 46, 1959 (B<sub>1</sub>H<sub>6</sub>), Stock, Friederici, and Priess, Ber., 1913, 46, 3353 (B<sub>18</sub>H<sub>14</sub>); Stock, Zeitsch. Elektrochem., 1913, 19, 779.

(5) R<sub>10</sub>H<sub>14</sub>. Obtained like (1); also by heating (1).
(6) Non-volatile solid, soluble in CS, Obtained like (1).

(7) Non-volatile yellow solid, insoluble in CS,. Obtained like (1), and by heating (6)

(8) Difficultly volatile liquid, obtained by heating (6).

(9) Colourless, non-volatile solid, insoluble in CS<sub>2</sub>. Obtained by heating (1).

(10) Brown hydride or hydrides, resembling boron and poor in hydrogen content Obtained by heating (7)

The starting-point in the preparation of the boron hydrides is crude magnesium boude, made by mixing 1 part of very finely powdered boron sesqui oxide with 3 parts of magnesium powder and rapidly heating the mixture, 10 gianis at a time, in a thin sheet-iion crucible in a stream of hydrogen The product, quickly cooled in hydrogen, is finely powdered, sifted, and slowly dropped into 15 per cent hydrochloric acid at 50° to 80°, a slow stream of hydrogen being passed through the apparatus The evolved gases, dried over calcium chloride and phosphoric anhydride, are passed through U-tubes cooled in liquid air , in them the boron hydrides B, H, o and

 $B_0H_1$ , solidify, together with a little carbon dioxide and silicon hydrides. The distillation of this crude product furnishes in order the following fractions (i.) silicon hydride, SiII, (ii) carbon dioxide, (iii.) silicon hydride, Si\_2H\_0, (iv.) boron hydride, B\_4II\_{10}, (v) boron hydride, B\_6II\_{12}, (vi.) residue of less volatile boron and silicon hydrides

The two hydrides B\_4II\_10 and B\_6II\_12 are obtained as follows -- At the temperature of liquid air the pressure above the solid is reduced to zero. Traces of hydrogen and silicon hydride are thus removed. The temperature is then raised to -80°, when the solid melts. The pressure is quickly lowered to a few millimetres, and the evolved gas rejected as long as it contains silicon hydride. The gas then evolved at -80° and 3 mm, consists of the hydride B4H10, and, unless the room temperature is above 20°, partly condenses in the mercury pump is distilled at  $-40^\circ$  until the pressure falls to  $1.5~\mathrm{mm}$ , the temperature raised to 0°, and distillation continued until the pressure is only 10 mm. All the hydride  $B_4H_{10}$  has then been removed, and the residue, on further distillation, gives another hydride, B, H,, the pressure falling to less than 5 mm.

From 200 grams of magnesium boride, 100 c.c. (at N.T P.) of pure hydride

B<sub>4</sub>H<sub>10</sub> and 60 milligrams of hydride B<sub>6</sub>H<sub>12</sub> may be obtained.<sup>1</sup>

Tetraborodecahydride or borobutane, B<sub>1</sub>H<sub>10</sub>, is a colourless, very volatile liquid which boils at 16° to 17° C, and freezes at about -112° C. The vapour density is 270 (II = 1), in accordance with the formula given. The vapour pressures at various temperatures are as follows -

The hydride has a very disagreeable odour A few bubbles when inhaled affect respiration and cause headache It is extremely unstable, decomposing in a few hours at ordinary temperatures into a series of other hydrides. It is decomposed by electric sparks, and ignites spontaneously in air or oxygen, burning with a green flame . Nitric acid oxidises it with explosive violence.

<sup>&</sup>lt;sup>1</sup> Stock and Massenez, loc. cit. For a number of improvements in the preparation, see Stock, Kuss, and Priess, Ber., 1914, 47, 3115 VOL. IV.

It is slowly decomposed by water, the final result being expressed by the equation :—

$$B_4H_{10} + 12H_2O - 4H_3BO_3 + 11H_2$$

Aqueous sodium hydroxide iapidly and completely absorbs the hydride, producing a hypoborate (p. 28), but hydrogen is slowly evolved, and the net result may be expressed thus—

$$B_1H_{10} + 4NaOH + 4H_2O - 4NaBO_2 + 11H_2$$
.

The hydride is decomposed by alcohol and reacts with aminoma, but its solution in benzene is very stable towards oxygen.

Hexaborododecahydride or borohexylene, B<sub>0</sub>U<sub>19</sub>, is a colourless liquid, the vapour density of which is in agreement with the formula given. The vapour pressures at various temperatures are as follows.—

The hydride has a highly disagreeable odom, and ignites spontaneously in the air. It is decomposed by water and alkalies more readily than the preceding hydride:— $^1$ 

$$\begin{array}{cc} {\rm B_{\rm d}H_{12} + 18H_2O} & {\rm 6H_3BO_3 + 15H_2} \\ {\rm B_{\rm o}H_{12} + 6NaOH + 6H_2O} & {\rm 6NaBO_2 + 15H_2} \end{array}$$

Boron trihydride or boro-ethane,  $B_1 H_{\nu}$ —It has been mentioned that the compound  $B_1 H_{10}$  is very unstable. When kept over mercury at the ordinary temperature it decomposes with the formation of a gaseous hydride  $B_2 H_{\theta}$  and hydrogen. The rate of decomposition is greatly augmented by ultraviolet light, but not appreciably by sunlight. The change is practically complete at 100° after one hour.

Boron trihydride is a colourless gas possessing a repulsive odour which recalls that of  $B_1H_{10}$ , and also that of hydrogen sulphide—Its vapour density corresponds with the molecular formula  $B_2H_0$ —The hydride melts at  $-169^\circ$  and boils at  $-87^\circ$  C. The vapour pressure of the liquid at various temperatures is as follows.—

Boron tilhydride is fairly stable, but it slowly decomposes even at the ordinary temperature. When heated with a fice flame, solid, but not liquid, hydrides are produced, when spacked, boron is deposited and non-volatile hydrides of unpleasant odour are produced. In the presence of air it gives rise to solids containing both boron and oxygen, and when exploded with oxygen it reacts thus:—

$$B_2H_6 + 3O_2 \cdot B_2O_3 + 3H_2O$$
.

It dissolves in carbon disulphide, and the solution decolorises bromine; it does not react with dry hydrogen chloride, but combines readily with dry ammonia.

Boron trihydride reacts with water to give boric acid and hydrogen:—

$$B_2H_6 + 6H_2O = 2H_3BO_3 + 6H_2$$

Like the hydride B<sub>1</sub>H<sub>10</sub>, it gives a hypoborate with alkali hydroxide, but some hydrogen is evolved immediately the gas and the hydroxide come into

<sup>1</sup> Stock and Massenez, loc. cit.

contact. The hypoborate slowly decomposes, and the final result may be expressed thus:-

$$B_2H_6 + 6NaOH = 2Na_3BO_3 + 6H_2$$

Decaborotetra decahydride,  $\rm B_{10}H_{14}$  —When the hydride  $\rm B_4H_{10}$  is heated

for four or five hours to 100°, or when the hydride B<sub>3</sub>H<sub>6</sub> is heated for forty-eight hours to 115° to 120′, a volatile solid hydride, B<sub>10</sub>H<sub>10</sub> is produced.

This hydride is a colourless solid of peculiar odour resembling that of osmium tetrovide. It sublimes in vicuo at 60° to 80°, producing beautiful long. needles of density 0.94. It melts at 99.5 to a colourless liquid that does not decompose below 200° The molecular weight in benzene solution and the percentage composition are in harmony with the molecular formula  $B_{10} H_{14} \cdot$ 

The hydride is soluble in alcohol, ether, benzene, and carbon disulphide. It is not decomposed by air or water, is little affected by mitric acid, but is oxidised by potassium permanganate. It dissolves in alkalies, forming a yellow solution

When prepared from B<sub>1</sub>H<sub>10</sub> or B<sub>2</sub>H<sub>6</sub> by heating, the hydride B<sub>10</sub>H<sub>14</sub> is accompanied by two other non-volatile solid boron hydrides, which are also produced when B<sub>10</sub>H<sub>11</sub> is heated (i.) a yellow hydride, probably (B<sub>5</sub>H<sub>4</sub>)<sub>2</sub>, insoluble in carbon disulphide and decomposed by water, and (ii) a colourless hydride, probably B<sub>12</sub>H<sub>x</sub>, soluble in carbon disulphide, not decomposed by water, and converted into (1) at 150'.2

Borides .-- The binary compounds of boron with carbon and silicon are described in this chapter as boron carbide and boron silicides. The binary compounds of boron with other, more electropositive elements, are called borides, and are described under the headings of the other elements.

### BORON AND THE FLUORINE GROUP.

Boron forms halogen compounds of the type BX<sub>8</sub>. The fluoride, schloride, and bromide can be prepared by the direct union of their elements, but not the iodide. The chloride and bromide can also be prepared by heating to redness in a stream of either chlorine or bromine an intimate mixture of boron sesqui-oxide and carbon -

$$B_2O_3 + 3O + 3X_2 = 2BX_3 + 3CO$$
.

The boion halides have none of the properties of salts. At the ordinary temperature the fluoride is a gas, the liquid chloride boils at 12 5° C., the bromide is a volatile liquid, and the iodide a solid of low melting-point. Each is rapidly decomposed by water, the change being expressed, except for the fluoride, by the (irreversible) equation -

$$\mathrm{BX_3} + 3\mathrm{H_2O} = \mathrm{H_3BO_3} + 3\mathrm{HX}.$$

Besson has described the bromo-iodides of boron, BBr 1 and BBr 12. They are colourless liquids which boil at 125° and 180° respectively. A mixture

Stock and Friederici, loc cit., Stock, Friederici, and Priess, loc. cit.
 Stock, Friederici, and Priess, loc. cit., Stock, Zeitsch. Elektrochem., 1913, 19, 779.
 Moissan, Compt. rend., 1904, 139, 364.
 Herzelius, Poyg. Annalen, 1824, 2, 147.
 Wohler and Deville, And Chim. Phys., 1858, [iii], 52, 63.
 These syntheses have only been effected with "amorphous boron"
 Dumas, Ann. Chim. Phys., 1826, [iii.], 31, 436.
 Poggalo, Compt. rend., 1846, 22, 127.
 Besson, Compt. rend., 1891, 112, 1001.

of these compounds and boron tri-iodide is obtained when hydrogen iodide acts upon boron tribromide at a high temperature.

Two oxychlorides of boron, BOCl and BOCl, have been described, but it is doubtful whether they really exist. The first is stated by Gustavson to be obtained as a white, gelatinous solid when boron sesqui-oxide and boron trichloride are heated together at  $150^\circ$  in a scaled tube.

Boron trifluoride, BF2, was originally prepared by heating a mixture of boron oxide and calcium fluoride to a white heat in an iion tube.2 It may be prepared by heating a mixture of boron throude (1 pt ) and calcium fluoride (2 pts.) with concentrated sulphuric acid,3 or better, by mixing 100 parts of potassium borofluoride with 15 to 20 parts of powdered boron sesqui-oxide and heating with concentrated sulphunic acid 4. The gas may be collected over mercury.

Boron trifluoride is a very stable, colourless gas of suffocating odour. Its density corresponds to that required for the simple formula BF3 When purified from hydrogen fluoride by passage over sodium fluoride, it melts at and boils at -101°.5 The figuefaction of the fluoride was first effected by Faraday.6 Boron trifluoride combines with its own volume of ammonia, producing a white, opaque solid,  $BF_3$  NH, which can be sublimed without decomposition. Liquids of the composition  $BF_3$  2NH, and  $BF_3$  3NH, may be obtained if more animonia is employed (J. Davy), but their individuality is doubtful. With dry phosphine at - 50°, boron fluoride unites to form a white, unstable solid of the formula 2BF<sub>3</sub> PH<sub>3</sub> 7 It also combines with the oxides of nitrogen.8

Boron trifluoride is rapidly absorbed by water, 1057 volumes being absorbed at 0° C 9 The products of the reaction are bone acid and hydrofluoboric acid .-

$$4(BF_3) + 3H_2O + Aq. = 3HBF_4Aq + H_3BO_3Aq. + 98.04 \text{ Cals.}^{10}$$

Owing to the readiness with which boron trifluoride reacts with water, it has been proposed as a dehydrating agent in organic chemistry 11

Boron trifluoride appears to combine with hydrogen fluoride in two proportions, giving rise to BF, HF or HBF, and BF, 3HF Little or nothing is known of the latter beyond the fact that it is a colourless liquid of normal vapour density.12 The former is a well-known acid, hydrofluoboric acid

Hydrofluoboric acid, HBF, was discovered by Berzelius 13 - It is obtained in aqueous solution when boron trifluoride is passed into water until the liquid reacts strongly acid, and the solution is then cooled The boric acid simul-

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    Gustavson, Ber., 1870, 3, 426, J. Russ, Chem. Soc., 1888, 20, 621. R. Loienz, Annalen, 1888, 247, 226; Counclei, Ber., 1878, 11, 1106; J. prakt. Chem., 1878, [ii], 18, 399;
    Michaelis and Becker, Jier., 1881, 14, 91;
    Gay-Lussac and Thenard, Ann. Chim. Phys., 1908, 69, 201, Rechecker physico-
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<sup>&</sup>lt;sup>a</sup> Gay-Lussac and Thenard, Ann. Chim Phys., 1908, 69, 201, Recherches physicochimiques, vol. ii. p. 38
<sup>a</sup> J. Davy, Phil. Trans., 1812, 102, 365, Berzelius, Pony. Annalen, 1824, 2, 113; Moissan, Compt. vend., 1904, 139, 711.
<sup>4</sup> Schiff, Annalen Suppl., 1867, 5, 172.
<sup>5</sup> Moissan, Lec. c.t., and Ann. Chim. Phys., 1906, [viii], 8, 84.
<sup>6</sup> Faraday, Phil. Trans., 1845, 135, 155.
<sup>7</sup> Besson, Compt. rend., 1890, 110, 80
<sup>8</sup> Kuhlmann, Annalen, 1841, 39, 319.
<sup>9</sup> Bassrow, Compt. rend., 1871, 78, 1698.
<sup>10</sup> Hemmerl, Compt. rend., 1880, 90, 312.
<sup>11</sup> Landolph, Compt. rend., 1877, 85, 39.
<sup>12</sup> Landolph, Compt. rend., 1878, 86, 639, 601, 671, 1463, 1879, 89, 173, Ber., 1879, 12, 1583.
<sup>12</sup> Landolph, Compt. rend., 1878, 86, 601.
<sup>13</sup> Berzelius, Pogg. Annalen, 1814, 2, 113.

<sup>18</sup> Berzelius, Pogg. Annalet, 1814, 2, 113.

taneously formed is said to separate out as metaboric acid. An aqueous solution may also be prepared by dissolving bone acid in a cold, dilute, aqueous solution of hydrogen fluoride

In aqueous solution, hydrofluobone and reacts as a monobasic complex acid, the ions of which are H and  $BF_4'$ . Apparently there is also a slight secondary dissociation of the ion  $BF_4'$  into  $BF_4$  and F', the  $BF_3$  partly hydrolysing to hydrofluoric and boric acids. The diluted acid does not attack glass, when the solution is concentrated, however, it does so, since hydrogen fluoride is produced The acid is poisonous

The salts of hydrofluobone acid are called borofluorides. They may be prepared (1.) by neutralising the acid with metallic hydroxides, oxides, or carbonates, (ii ) by the action of the so-called fluobone acid (p. 21) on metallic fluorides, and (iii.) by the action of a metallic fluoride and hydrofluoric acid, or an acid fluoride, on borie acid. In the last case the curious phenomenon may be observed of a mixture of two acid liquids producing an alkaline solution, eg. -

$$H_3BO_3 + 2NaHF_2 = NaBF_1 + NaOH + 2H_2O$$
.

Most borofluorides are soluble in water. The insolubility of the potassium salt in 50 per cent alcohol may be utilised for separating potassium from sodium and magnesium. When heated to redness, borofluorides yield boron trifluoride and a metallic fluoride, with concentrated sulphuric acid, boron trifluoride and hydrogen fluoride are evolved. A mixture of fluoride and borate is produced by fusing a borofluoride with an alkali carbonate.2

Potassium borofluoride, KBF, may be prepared as follows -Boric acid (1 part) and calcium fluoride (2 5 parts) are heated with boiling sulphuric acid (5.25 parts) The liquid is cooled and filtered, and a solution of a potassium salt added Potassium borofluoride is precipitated; it is recrystallised from hot water 3 Thus obtained, it forms anhydrous, shining, six-sided, prismatic crystals of density 2.5, and is orthorhombic (a: b:c=0.7898:1:1:2830), being isomorphous with the rubidium salt, RbBF<sub>4</sub> (a:b:c=0.8067·1·1·2918, density=2.820). When prepared from hydrofluoric acid, boric acid, and potassium carbonate, however, it forms a gelatinous mass which is converted at 100" into crystals belonging to the cubic system and showing the faces of the octahedron and dodecahedron.4

The following borofluorides are also known (Berzelius) -

LiBF, NaBF,	CsBF <sub>1</sub> <sup>6</sup> (NH <sub>1</sub> )BF <sub>1</sub> <sup>7</sup>	Ca(BF <sub>1</sub> ) <sub>2</sub> + H <sub>2</sub> O Ba(BF <sub>1</sub> ) <sub>2</sub> , H <sub>2</sub> O	Pb(BF <sub>1</sub> ) <sub>2</sub> Cu(BF <sub>4</sub> ) <sub>2</sub>
KBF,	$Mg(BF_1)_2 \times H_2O$	$Al(BF_4)_3$ x $II O$	Ou(13F <sub>4</sub> / <sub>2</sub>
RbBF.	Zn(BF.).	Y(BF.), H.O	

Fluoboric acid.--When boron trifluoride in sufficiently large amount . is passed into cold water, a colourless, syrupy liquid is produced. A similar liquid is obtained when as much boric acid as possible is dissolved in a cold, concentrated solution of hydrofluoric acid and the solution concentrated on

<sup>&</sup>lt;sup>1</sup> Mathers, Stewart, Housemann, and Lee, J. Amer. Chem. Soc., 1915, 37, 1515.

<sup>2</sup> Marignac, Zeitsch. anal. Chem., 1862, 1, 405.

<sup>3</sup> Stolba, Chem. Zentr., 1872, p. 395; 1875, p. 403; 1876, p. 703.

Montemartini (and Brugnatelli). Altis. R. Accad., Lincei, 1894, [v.], 3, 1. 339; Gazzetta, 1894, 24, 1. 478; Zeitsch. Kryst. Man., 1894, 26, 198.

<sup>5</sup> Zambonni, Zeitsch. Kryst. Min., 1905, 41, 57.

<sup>6</sup> Godeffroy, Ber., 1876, 9, 1867.

<sup>7</sup> Stolba, Chem. Zentr., 1890, 1. 211. •

the water-bath, boiled until it evolves white fumes, and cooled over concentrated sulphuric acid.1

The liquid thus obtained, of density 1 584, has a composition in agreement with the formula II, B,O, 6HF. It is decomposed by excess of water with the precipitation of boric acid, an aqueous solution of hydrofluoboric acid being produced Berzelius regarded the liquid as a definite acid, of which he could form the alkali salts M2B2O46MF H2O by neutralisation with alkali. This view, however, is in all probability erroncous, although the evidence against the individuality of the liquid is not as conclusive as could be desired. Distillation of the liquid is accompanied by rise of boiling-point, and the successive hquid fractions differ in density and composition, whilst initially boron trifluoride is evolved. The "salts, when recrystallised, yield first MF and subsequently a mixture of MF and  $M_2B_2O_4$ , while the "acid" in aqueous solution gives with silver nitrate a precipitate of silver metaborate mixed with silver oxide According to Basarow, the liquid is merely a mixture of metaboric, hydrofluoboric, and hydrofluoric acids.2

Two other fluoboric acids have been stated to exist, namely, H<sub>1</sub>B<sub>2</sub>O<sub>7</sub>.3HF and H<sub>4</sub>B<sub>2</sub>O<sub>9</sub> 2HF, but little is known concerning them <sup>3</sup> A physico-chemical study of mixtures of boric acid, hydrofinoric acid, and potassium fluoride has been made by Abegg, Fox, and Herz, but no definite conclusions could be

Although no fluoboric acids are definitely known, two compounds have been prepared which may be looked upon as salts of such acids. These are B<sub>2</sub>O<sub>8</sub>.2KF and KBO<sub>2</sub> KF, to which the constitutions

$$F$$
 BOB  $F$  and  $F$   $F$   $O.K$ 

have been assigned. The former is made by fusing 7 parts of boron sesquioxide with 12 parts of potassium fluoride, allowing the melt to cool slowly, and extracting soluble impurities with alcohol; the latter, by fasing the former with the requisite amount of potassium carbonate. The compounds dissolve without decomposition in a little water, but much water decomposes them.5

Perfluoboric acid -This acid is not known, but several perfluoborates have been prepared. Potassium fluohyperborate,  $K_1B_1F_4O_{11}.H_2O$ , is prepared by dissolving the compound  $B_2O_32KF$  in cold water, adding a slightly alkaline solution of hydrogen peroxide, then adding alcohol and stirring. The dry salt is a white, crystalline solid. The aqueous solution is alkaline and slowly evolves oxygen, dilute sulphuric acid added to the solution liberates hydrogen peroxide Concentrated sulphuric acid decomposes

the substance, ozonised oxygen being evolved.

The potassium salt, K<sub>2</sub>B<sub>2</sub>F<sub>2</sub>O<sub>0</sub>, H<sub>2</sub>O or O<sub>2</sub>(BF.O.OK)<sub>2</sub> H<sub>2</sub>O, and the ammonium salt,  $(NH_1)_2B_2F_2O_6.3H_2O$ , have also been obtained by

<sup>&</sup>lt;sup>1</sup> Gay Lussac and Thenaid, Ann. Chim. Phys., 1809, 69, 201; see also Berzehus, Pogg. Annalen, 1843, 58, 503, 1843, 59, 644.

<sup>2</sup> Basarow, Compt. rend., 1374, 78, 1698; Ber., 1874, 7, 1121.

<sup>3</sup> Landolph, Ber., 1879, 12, 1583.

<sup>4</sup> Abegg, Fox, and Herz, Zeitsch. anorg Chem., 1903, 35, 129.

<sup>5</sup> Schiff and Sestini, Annalen, 1885, 228, 83.

<sup>6</sup> Melkoff and Lordkipanidze, Ber., 1899, 32, 3349, 3510; see also Petrenko, J. Russ. Phys. Chem. Soc., 1902, 34, 37.

Boron subchloride (?).—When a mixture of boron trichloride vapour and hydrogen at the ordinary temperature is subjected to the action of the silent electric discharge, a pale yellow, transparent substance is produced which contains fewer than three atoms of chlorine per atom of boron. It is rapidly decomposed by water, hydrogen being evolved and a solution obtained

which possesses marked reducing properties.1

Boron trichloride, BCl<sub>9</sub>, is best prepared by either of the two methods already mentioned (p. 19). For the synthesis, impute boron (prepared by heating the oxide with magnesium and boiling the resulting mass with hydrochloric acid) is employed. It is heated to dull redness in a stream of dry hydrogen, allowed to cool in the gas, and, after displacing the hydrogen with chlorine, heated in chlorine to redness. The product is strongly cooled and protected from moisture, shaken with mercury or silver powder to remove chlorine, and fractionally distilled to eliminate hydrogen chloride and silicon chloride <sup>2</sup> Boron trichloride is also produced when boron sesqui-oxide is heated for some days at 150° with phosphorus pentachloride <sup>3</sup> Boron trichloride is a colourless liquid of high refractive index and normal

vapour density (Wohler and Deville)  $^{+}$  At 0°  $^{0}$  its density is 143386;  $^{1}$  its coefficient of expansion is large. It melts at  $-107^{\circ}$  C and boils at 125° C.; its vapour pressure is as follows.

°C. nim	'C   m	n " ('.	mm	0.0	mm.	" ('.	, mm
-80   4 75   6 70   9 -65   13	-55 -50	8 40 6 35 7 - 30 1 25	67 89 116 151	- 20 - 15 - 10 - 5	197 251 311 387	0 5 10 12 4	477 579 695 758

One gram-molecule of the trichloride, when decomposed by water, produces 79 2 Cals 9 Boron trichloride is without action on sodium below 150°, zinc below 200°, and sulphur below 250°, but it reacts with sulphur trioxide at 120°, producing sulphuryl chloride and boron sesqui oxide (Gustavson).

When boron trichloride vapour is passed into liquid ammonia, excess of which is then removed at  $-23^{\circ}$ , 15 molecules of ammonia are fixed for each molecule of the chloride used. The products are boron amide and an ammonia addition-product of ammonium chloride --

$$BCl_3 + 15NH_3 - B(NH_2)_3 + 3NH_4Cl.3NH_3$$

If the ammonia be removed at 0°, only 6 molecules are fixed, since the dissociation pressure of NH  $_2$ Cl 3NH  $_4$  exceeds 760 mm at 0°. When the reaction takes place above 0° C, boron mude, B,(NII)30 is produced :-

$$2B(NH_2)_1 B_2(NH)_3 + 3NH_3$$
.

<sup>1</sup> II V A. Birscoe; private communication,
2 Schnitzler, Dingl. Folgl. J., 1874, 211, 485; Wohler and Deville, loc., cil.; Stock and
Priess, Ber., 1914, 47, 3109.
3 Gustavson, Ber., 1870, 3, 426; 1871, 4, 975.
4 Ghita, Zeitsch physikal Chem., 1893, 12, 768, Garzetta, 1893, 23, 1, 452.
5 Stock and Priess, loc. cit., cf. Regnault, Mem. Acad. Sci., 1862, [ii.], 26, 479, 658.
6 Troost and Hautefeuille, Ann. Chim. Phys., 1876, [v.], 9, 70.
7 For other reactions see Troost and Hautefeuille, Compt. rend., 1872, 75, 1819; Ann.
Chim. Phys., 1876, [v.], 7, 476.
5 Joannis, Compt. rend., 1902, 135, 1106.

According to Besson,1 boron trichloride forms a white, solid compound, BCl, PH, with phosphine. The following double compounds have also been described, besides various addition-compounds with organic substances:

BCl<sub>3</sub>.NOCl, BCl<sub>3</sub>.POCl<sub>3</sub>, BCl<sub>3</sub>.CNCl, and BCl<sub>3</sub>.HCN.<sup>2</sup>
When boron trichloride is introduced into the afterglow of active nitrogen (see Vol. VI.), a pale bluish-green glow is developed, easily distinguished from the green colour of the boric acid flame, and a white, amorphous solid that contains both boron and nitrogen (probably boron nitride) is deposited a

Boron tribromide, BBr3, is best prepared by either of the two methods already mentioned (p 19); its preparation and purification may be carried out as described for the chloride 4.5

Boron tribromide is a colourless, mobile liquid which boils at 90 5° C. 4, 5 and solidifies to a colourless mass which melts at  $-46^{\circ}$  C,  $^{5,6}$  Its density is 2 6499 at 0° C and 2 6175 at 16 5°.7 The vapour density is normal;  $^{4}$ the vapour pressure is as follows .- 5

°C.	mm.	° ر'.	mm,	° C	mm.	° C.	mın	° C.	mm
50	0·7	- 20	5 5	10	32	40	130	70	890
- 40	1 5	- 10	10 5	20	54	50	193	80	540
- 30	3	0	19	30	85	60	280	90	730

Boron tribromide reacts with ammonia at 0° to form boron imide.

$$2BBr_3 + 27NH_3 = B_2(NH)_3 + 6NH_4Br 3NH_3.8$$

With phosphine, boron tribromide forms a white, solid substance of the formula BBr, PH, 9 The corresponding arsenic compound BBr, AsII, prepared by the direct union of arsine and boron tribronide at -100, is a very unstable, white, amorphous solid, which, on keeping in the dark at 0°, slowly decomposes into boron tribromide, arsenic, and hydrogen 10 The following addition-compounds are also known BBr, PBr, BBi, PBR, 2BBr, PCl,

2BBr<sub>3</sub> PCl<sub>5</sub>, and 2BBr<sub>3</sub>, P<sub>2</sub>I<sub>4</sub> <sup>11</sup>
Boron tri-iodide, BI<sub>3</sub>, was first prepared by Mossan in 1891. He obtained it by three different methods. (i.) by acting upon boron trichloride vapour with hydrogen iodide at a high temperature, (ii) by the action of iodine vapour on "amorphous boron" at 700° to 800°, and (iii.) by the action of hydrogen iodide on "amorphous boron" at a red heat. The third process is the best to employ. The "boron" to be used must be obtained by Wohler and Deville's method (p 7), washed with hydrochloric acid, and dried at 200° in a current of hydrogen. It is heated in a Bohemian glass tube in a current of hydrogen iodide (dried over calcium iodide) to a

<sup>&</sup>lt;sup>1</sup> Besson, Compt rend., 1890, 110, 516.

<sup>1</sup> Besson, Compt rend., 1890, 110, 516.
2 Genther, J. prakt. Chem, 1873, [I.], 8, 357; Gustavson, Bcr., 1871, 4, 975, Maitius, Amalen, 1859, 109, 80; Gautier, Compt. rend., 1866, 63, 920.
3 Javons, Proc. Roy. Soc., 1915, A, 91, 120.
4 Wohler and Deville, loc. c.t.
5 Stock and Kuss, Ber, 1914, 47, 3113.
6 H. Gautier, Ann. Chim. Phys., 1899, [vii.], 18, 374.
7 Ghira, loc. cit.; Hoskyns-Abrahall, Trans. Chem. Soc., 1892, 61, 650.
9 Joannis, Compt. rend., 1904, 139, 364; cf. Besson, ibid., 1892, 114, 542.
9 Besson, abid., 1891, 113, 78.
18 Stock, Ber., 1901, 34, 949.
14 Tarible, Compt. rend., 1893, 116, 1521; 1901, 132, 83, 204.
18 Moissan, Compt. rend., 1891, 112, 717; cf. Besson, ibid., 1891, 112, 1001.

temperature just below that at which the glass softens. The crystalline product is dissolved in carbon disulphide, shaken with mercury to remove iodine, and the boron iodide recovered by allowing the carbon disulphide to

Boron tri-iodide crystallises in colourless, transparent, nacreous plates which are very hygro-copic and are easily changed by light. It melts at 43° and boils without decomposition at 210". It is soluble in carbon

disulphide, carbon tetrachloride, benzene, and other organic media.

Boron tri-rodide is not attacked by hydrogen It is decomposed by sodium and by magnesium at a red heat, but is unaffected by silver at 500° and by sodium at 210°. The iodine buins in oxygen, is attacked by phosphorus at the ordinary temperature, and by sulphur when gently warmed

According to Besson, boron tri-iodide forms the compound BI 5NH 3 with ammonia, and also unites with phosphine.

#### BORON AND THE OXYGEN GROUP.

Oxides of Boron -- At least four oxides of boron appear to exist, viz.  $B_4O_3$ ,  $B_2O_3$ ,  $B_4O_3$ , and  $B_2O_3$ . Of these, the sesqui-oxide,  $B_2O_3$ , has been long known, and is a well-defined acidic oxide. The dioxide,  $B_2O_2$ , is also acidic.

Tetraboron trioxide, B<sub>1</sub>O<sub>3</sub> -When boson dioxide is prepared by method (11.) described below, it is found that about five per cent of the product consists of a substance insoluble in hydrochloric acid or sodium hydroxide but readily soluble in intric acid with conversion into boric acid. This insoluble product, when dried at 100, appears to consist of a hydrated oxide of the composition B<sub>1</sub>O<sub>3</sub> 2H<sub>2</sub>O<sup>2</sup>
Boron dioxide, B<sub>2</sub>O<sub>2</sub>.—This oxide may be prepared as follows. Crude

magnesium boulde, made as described below, is decomposed by cold water; the filtered solution is then treated by either of the following processes: (1.) the solution is evaporated to dryness in vacuo and the residue heated; (ii.) the solution is treated with ammonia, filtered from magnesium hydroxide, and evaporated to dryness in vacuo and the residue heated. In method (1.) part of the boron dioxide obtained is present in combination with magnesia (as a magnesium borite); in method (ii ) the dioxide is obtained practically free from magnesia but contaminated with about five per cent of the lower oxide B<sub>4</sub>O<sub>8</sub>.2H<sub>2</sub>O.

Boron dioxide is soluble in water Its aqueous solution can be oxidised to boric acid by evaporation in air or by heating with intric acid, but is not affected by rodine. Freezing-point measurements indicate that the molecular formula of the oxide is B2O2. The oxide apparently combines with water to form an acid, and the aqueous solution dissolves an amount of freshly precipitated magnesium hydroxide corresponding to the production of a

magnesium borite, MgO 2B,O, 2 Tetraboron pentoxide, B4O5, is prepared by treating the compound

Mg8B2(OH), with strong ammonia for several days in an atmosphere of hydrogen, filtering, evaporating the filtrate to dryness in vacuo, and heating the residue. Thus prepared, it forms a pale brown solid, infusible at the

Besson, Compt. rend., 1892, 114, 542 Travers, Ray, and Gupta, J. Indian Inst Sci., 1914, I, 1. 1, and private communication from Dr Travers

softening point of Jena glass. When allowed to remain in vacuo in contact with water it dissolves, forming a golden-yellow solution that oxidises rapidly in the air; a small flocculent residue, apparently consisting of B<sub>1</sub>O<sub>2</sub> 2H<sub>2</sub>O, is also obtained. It is possible that the residue is produced by the following reaction: -- 1

$$3B_1O_5 - B_1O_3 + 4B_2O_3$$

Borohydrates.-When one part of boron sesqui-oxide is heated with 21 parts of magnesium powder for 15 minutes at a red heat in a rapid stream of hydrogen, the main products are magnesia and magnesium boride,  $Mg_8B_2 :-- ^2$ 

$$6Mg + B_2O_3 = 3MgO + Mg_3B_2$$
.

It appears to be essential to the success of the experiment that the mixture should gently dellagrate for about five minutes after the heating commences.

The crude mixture from the foregoing reaction evolves hydrogen when treated with water. It has been shown by Travers, Ray, and Gupta's that the magnesium boride is decomposed as follows

$$Mg_3B_2 + 6H_2O = Mg_3B_2(OH)_6 + 3H_2$$

The product is an almost white powder, insoluble in water, it is the magnesium derivative of a compound B<sub>2</sub>(OH<sub>2</sub>)<sub>6</sub>, which, from analogy with the carbohydrates, may be termed a borohydrate.

The solution obtained by treating the boride with water is found to contain small quantities of substances which exhibit the remarkable property of evolving hydrogen, with brisk effervescence, when acidited. These substances are also borohydrates or allied compounds, and, like the hydroborons obtained from the magnesium boride by the action of acids, appear to be the products of certain unknown side reactions. The amounts of these substances obtained are very small in comparison with the amount of bonde required for their production.

The solution is usually yellow, owing to the presence of colloidal boron. It decomposes slowly at the ordinary temperature, hydrogen being evolved In the presence of platinum black the rate of decomposition is greatly accelerated. The solution precipitates silver and incremy from their salts immediately; with copper salts either copper hydride or amorphous boron appears to be precipitated, according to circumstances. When acidified, the solution evolves hydrogen; the liquid thus obtained decolorises iodine.

A careful, quantitative study of the properties of the solution has led Travers, Ray, and Gupta 4 to the conclusion that the solution contains two substances of the formulæ  $H_0B_2O_2$  and  $H_0B_2O_3Mg$ , the latter being the magnesium derivative of a compound  $H_nB_2O_2$ . It is suggested that in these compounds boron has a valency of five; and if, as seems highly probable, the evolution of hydrogen takes place by the elimination of pairs of hydrogen

<sup>&</sup>lt;sup>1</sup> Travers and Ray, J. Indian Inst. Sci., 1914, x, x, 97.

<sup>2</sup> See Ray, Trans. Chem. Soc., 1914, 105, 2162; Travers, Ray, and Gupta, J. Indian Inst. Sci., 1914, x, i. 1.

<sup>3</sup> Travers, Ray, and Gupta, loc. cit.

<sup>4</sup> Travers, Ray, and Gupta, loc. cit.; cf. Travers and Ray, Proc. Roy. Soc., 1912, A,

atoms attached to adjacent boron atoms, the properties of these compounds can be explained by assigning to them the formule  $-\!-\!$ 

The products formed on treatment with and arise in the following manner —

$$\begin{aligned} \text{(a)} \quad & \frac{\text{II}_2\text{B OH}}{\text{II}_2\text{B OH}} = \frac{\text{B OH}}{\text{B OH}} + 2\text{II}_2 \\ \text{(b)} \quad & \frac{\text{IIB}(\text{OH})_3}{\text{IIBH}_3} = \frac{\text{B}(\text{OH})_3}{\text{BH}_3} + \text{H}_2 \end{aligned}$$

The compound HO.B. BOH, which is analogous to hypomtrous acid in its structure, and the compound  $BH_3$ . B(OH), are oxidised by iodine to boron dioxide,  $B_2O_2$ :—

(iv.) 
$$\frac{B.OH}{B.OH} + I_2 = \frac{B.O}{B.O} + 2HI$$
  
(iv.)  $\frac{BH_3}{B(OH)_1} + 2I_2 = \frac{B.O}{B.O} + 4HI + H_2O$ 

When the solution obtained by the action of water on magnesium bonds is treated with ammonia, magnesium hydroxide is precipitated. In the conversion of the magnesium derivative  $\Pi_b B_2 O_3 Mg$  into the ammonium compound, however, intra-molecular change apparently occurs, thus —

$$\frac{\mathrm{HB}(\mathrm{OH})_3}{\mathrm{HBH}_4} \rightarrow \frac{\mathrm{H_2B}(\mathrm{OH})_2}{\mathrm{H_2BH}(\mathrm{OH})},$$

for the new product, when acidified, evolves twice as much hydrogen as the initial  $-\!\!\!-\!\!\!-$ 

$$\frac{H_2B(OH)_2}{H_3BH(OH)} = \frac{B(OH)_2}{BH(OH)} + 2H_2,$$

yielding a product which is oxidised by iodine to boron dioxide, B2O2:--

$$\frac{\mathrm{B}(\mathrm{OH})_2}{\mathrm{BH}(\mathrm{OH})} + \mathbf{I}_2 = \frac{\mathrm{B} : \mathrm{O}}{\dot{\mathrm{B}} : \mathrm{O}} + 2\mathrm{HI} + \mathbf{I}\mathbf{I}_2\mathrm{O}.$$

The magnesium derivative Mg<sub>3</sub>B<sub>2</sub>(OH)<sub>6</sub>, already mentioned, undergoes decomposition when treated with strong ammonia in an atmosphere of hydrogen. Travers and Ray, who have investigated the reaction, conclude that the soluble product of the change is a di-ammonium derivative of the compound—

When acidified, this compound loses hydrogen, thus -

$$\begin{array}{l} \mathrm{BH_2(OH)_2\text{-}BH(OH):BH(OH):BHI,(OH)_2} \\ \bullet = 2\mathrm{H_2} + \mathrm{BH(OH)_2} \\ \end{array} \text{B(OH):BH(OH)_2},$$

<sup>&</sup>lt;sup>1</sup> Travels and Ray, & Indian Inst. Sci., 1914, I, x. 97.

and the new product loses more hydrogen when treated with iodine:-

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\begin{split} \mathrm{BH}(\mathrm{OH})_2 : \mathrm{B}(\mathrm{OH}) : \mathrm{B}(\mathrm{OH}) : \mathrm{BH}(\mathrm{OH})_2 + \mathrm{I}_2 \\ &= 2 \mathrm{HI} + \mathrm{B}(\mathrm{OH})_2 \cdot \mathrm{B}(\mathrm{OH}) \; \mathrm{B}(\mathrm{OH}) \; \mathrm{B}(\mathrm{OH})_2 \cdot \mathrm{B}(\mathrm{OH}) \end{split}
```

By evaporating the ammoniacal solution in vacuo and heating the residue, the oxide  $B_4O_5$  is obtained (p. 25), thus —

$$\begin{aligned} BP_2(OH)_2 \cdot BH(OH) \cdot BH(OH) \cdot BH_2(OH)_2) &+ 2NH_3 \\ &= BO : BO \cdot BO \cdot BO + 5H_2 + H_2O + 2NH_3 \\ &- O \quad \_\_| \end{aligned}$$

**Hypoborates.**—Stock and Kuss¹ have shown that when either of the hydrides  $B_2H_6$  or  $B_4H_{10}$  dissolves in aqueous alkali hydroxide, the initial product is an alkali compound of the type MO.BH<sub>3</sub>, which they term an alkali hypoborate, eg

$$\begin{array}{l} {\rm B_2H_6} \ + 2{\rm KOH} = 2{\rm KOBH_3} + {\rm H_2} \\ {\rm B_1H_{10}} + 4{\rm KOH} = 4{\rm KOBH_3} + {\rm H_2}. \end{array}$$

Solutions of the alkali hypoborates are fauly stable at 0°, particularly when excess of alkali is present, and may be exposed to air. When boiled, however, decomposition takes place rapidly—

$$2KOBH_3 + 2H_2O = 2KBO_2 + 5H_2$$
.

Potassium hypoborate, KOBH, may be isolated by dissolving potassium hydroxide in one and a half times its weight of water and treating it with an excess of  $B_1H_{10}$  at 0°, the solid hypoborate is then obtained in colourless, glistening, octahedral crystals which may be dired in a high vacuum.

The compound is stable when dry—It is deliquescent, and its aqueous solution slowly decomposes at the ordinary temperature. Acids, even acetic acid, immediately decompose it—The aqueous solution is a powerful reducing agent, and gives precipitates with solutions of most salts, but insoluble hypotrates are never obtained. With copper sulphate, a precipitate of copper hydride, CuH<sub>2</sub>, is obtained; with nickel sulphate a very remarkable change takes place, nickel boride, Ni<sub>2</sub>B, being precipitated.

When potassium hypoborate is heated to 500°, potassium, hydrogen, and water are expelled. The water is evolved first, and is completely expelled at 200°; potassium begins to distil from the residue at 100°. The reaction may be expressed thus:—

$$5KOBH_3 = K_3B_3O_3 + 2K + 2H_2O + 11H.$$

The residue,  $K_3B_5O_3$ , is soluble in water, giving an alkaline solution the properties of which resemble those of a solution obtained by heating boric oxide with magnesium and extracting the product with water.

Sodium hypoborate, NaOBII<sub>3</sub>, resembles the potassium compound. The barium and magnesium hypoborates are only known in aqueous solution.

Boron sesqui-oxide, boron trioxide, or boric anhydride, B<sub>2</sub>O<sub>3</sub>, the best-defined of the oxides of boron at present known, is formed when boron burns in air or oxygen, and is prepared by heating boric and to redness in a platinum dish, water being eliminated and boron sesqui-oxide left behind.

5- -1

Boron sesqui-oxide is an amorphous, colourless, glassy solid, which has no definite melting-point, but which is decidedly liquid at  $557^{\circ}$ . It vaporises rapidly in vacuo at a bright red heat " Its density is 1.877 at 0°, 1.848 at 12°, and 1.699 at 80°, the coefficient of cubical expansion being 0.001308 Between 16° and 98' its specific heat is 0 2374.4 It is a very bad conductor of electricity.5

Boron sesqui-oxide is a very stable compound, but it can be reduced to boron by sodium, potassium, aluminium, and magnesium, as has been already mentioned (p 7). It is attacked by hydrofluoric acid and reacts with heated metallic fluorides, boron fluoride and crystalline metallic oxides being produced 6 Being only slightly volatile at a red heat, boion sesqui-oxide is capable of decomposing intrates, sulphates, etc., borates and volatile acids (or their decomposition products) being produced  $^7$ 

Boron sesqui-oxide is the anhydride of boric acid, into which it passes

when dissolved in water. The oxide is very hygroscopic

Boron sesqui-oxide combines with a number of other anhydrides to form what may be regarded as mixed anhydrides. The compound B<sub>2</sub>O<sub>3</sub>, P<sub>2</sub>O<sub>5</sub>, sometimes called borophosphoric acid or boron phosphate, was first prepared by Vogel 8 by adding crystalline boric acid to a boiling solution of phosphoric acid. According to Mylius and Meusser, when these two acids are heated together at  $80^\circ$  to  $100^\circ$  the product has the composition  $B_2O_vP_2O_g.2H_2O$ or (BO)H<sub>2</sub>PO<sub>4</sub> Borie and phosphoric acids also condense together in sulphune acid, acetic acid, or acetic anhydride as solvent, producing borophosphore acid This substance is a white powder, which, although readily hydrolysed by water when freshly prepared, is quite mert after it has been dried at 100" At a red heat, it becomes distinctly crystalline. mixed anhydrides SO, B,O, and 2SO,B,O, may be prepared by heating sulphuric and bone anhydrides together in sealed tubes at 115°-120° and 230 respectively, a mixture of both being obtained at intermediate temperatures to They are colourless, amorphous, hygroscopic solids which, when strongly heated, dissociate into sulphur and boron trioxides, and which are readily hydrolysed by water to sulphuric and boric acids. The compound (2B<sub>2</sub>O<sub>3</sub> 3SO<sub>3</sub> 3H<sub>2</sub>O or BHSO<sub>4</sub>), is produced by the interaction of boric and sulphuric acids; other compounds of the type xB<sub>2</sub>O<sub>1</sub> ySO<sub>3</sub> zH<sub>2</sub>O have been described, but their existence has not been confirmed 11

The existence of the preceding compounds is generally held to show that boron sesqui-oxide may act as a feeble base, in which case boron sesqui-oxide must be classed among the amphotene oxides 12

<sup>&</sup>lt;sup>1</sup> Carnelley, Trans. Chem. Soc., 1878, 33, 278.

<sup>2</sup> Trede and Bumbrauer, Zeitsch. anny. Chem., 1914, 87, 129.

<sup>3</sup> Ditto, Ann. Chim. Phys., 1878, [v.], 13, 67.

<sup>4</sup> Regnault, tibal., 1811, [in.], 1, 129.

<sup>5</sup> Lapschin and Tichanowitsch, Phil. Mag., 1861, [iv.], 22, 308; Bowgoin, Compt. rend., 1868, 67, 798.

<sup>6</sup> Deville and Caion, Compt. rend., 1858, 46, 764.

<sup>7</sup> Seo Tate, Quart. J. Chem. Soc., 1859, 12, 160.

<sup>8</sup> Vogel, Zeitsch. fur Chem., 1870, p. 125, Gustavson, Ber., 1871, 4, 975; Meyer, ibid., 1889, 22, 2919.

<sup>1889, 22, 2919.</sup> 

Mylus and Meusser, Ber 1901, 37, 397.
 Pictet and Karl, Bull. Soc. chim., 1908, [iv.], 3, 1114.
 D'Arcy, Trans. Chem. Soc., 1889, 55, 155, cf. Mer., J. makt. Chem., 1866, 99, 181;
 Schultz Sellack, Ber., 1871, 4, 16; Gugtavson. ibid., 1873, 6, 10.
 See Georgievič, J. prakt. Chem., 1888, [ii.], 38, 118.

Boric Acids.—Numerous boric acids are theoretically capable of being produced by the union of boron sesqui-oxide and water in different proportions, and the existence of various boric acids seems necessary in order to account for the types of borates known At the present time, however, the existence in the solid state of only two bone acids, orthoboric acid,  $H_3BO_3$  (i.e.  $B_2O_3$   $3H_2O$ ), and metaboric acid,  $11BO_2$  (i.e.  $B_2O_3$   $11_2O$ ), can be definitely affirmed. The latter is formed by heating the former to 100°-140°,1 and m aqueous solution it passes into the former, as is shown by molecular weight determinations. The existence of pyroboric acid,  $H_2B_1O_7$ , has been assumed by various chemists, but cannot be regarded as proved 2

Orthoboric acid, boric acid, or boracic acid, II, BO, -This acid was first prepared by Homberg m 1702 (see p 7) It may be readily prepared in the laboratory by treating a solution of borax (3 pts.) in hot water (12 pts) with sulphune and (1 pt.) On cooling, orthoboric acid crystallises out. It is recrystallised from hot water, dried, fused to expel traces of sulphuric acid, and again dissolved in hot water and recrystallised

Orthoboric acid is prepared on a commercial scale. Originally, all the orthoboric acid on the European markets came from Italy, and a large quantity of the acid is still derived from that country. It occurs in the sufficient, or jets of steam which issue from volcanic vents near Monte Rotondo, Lago Zolforeo, Sasso, and Larderello, in Tuscany Many borings have also been made in order to produce artificial sufficial. The sufficial are surrounded by brickwork basins, several of which are generally built on the side of a hill. Water from any convenient source is run into the uppermost basin and subjected to the action of the suffioni within it for a day. It is then run into the next lower basin, and so on, until the water contains about 2 per cent of boric acid. Next it is made to flow in a thin stream over a large sheet of cornugated lead, 2 metres wide and 125 metres long, placed on a slight incline and heated from below by the vapours from sufficing too poor in borie acid to be utilised for the extraction of the acid. Water may be evaporated in this way at the rate of 20,000 litres per day. The liquid that runs from the end of the lead sheet is further concentrated in leaden pans until the bone acid commences to crystallise out, the gypsum that is invariably deposited during the evaporation being removed from time to time. Crude Tuscan bone and contains 74 to 80 per cent of boric acid, 8 to 14 per cent, of ammonium and magnesium sulphates, 4.5 to 7 per cont of water, together with small quantities of gypsum, clay, sand, sulphur, organic matter, etc 3

The origin of the boric acid in the sufficient is not at present understood. It has been conjectured that the boric acid arises from a reaction between boron sulphide and water, on the other hand, it has been supposed to be produced from boron intride. According to Nasini, its source is the included tourmaline in the surrounding granite rocks, which yields boric acid when

Holt, Mem Manchester Phil Soc., 1911, 55, No. 10; cf. Merz, J. prakt Chem., 1866,
 17; Ebelmen and Bouqet, Ann Chem. Phins., 1846, [in.], 17, 63; Schaffgotsch, Poyy.
 Annalen, 1859, 107, 427; Bloxam, Quant. J. Chem. Soc., 1850, 12, 177.
 See the preceding references, and Nasmi and Ageno, Zeitsch. physikal. Chem., 1909,

<sup>&</sup>lt;sup>3</sup> Payen, Ann. Chim Phys, 1841, [ni.], I, 217, Wittstein, Repert. fur Pharm., 1840, 72, 175, Wohl, Dingl. polyt. J., 1866, 182, 173, Edmann, J. prakt. Chem., 1838, 13, 72.

heated in superheated steam 1 The gases issuing from the sufficient are radio-active, as also is the granitic rock from which they issue 2

Boric acid is prepared on a commercial scale from the various naturally occurring borates already described (p. 6). For this purpose they are sometimes dissolved in hot hydrochloric acid, and the boric acid, which crystallises out on cooling, recrystallised from water. Numerous other methods have been proposed. In one process, ulevite is ground to an impalpable powder, suspended in boiling water, and decomposed by passing sulphur dioxide into the liquid. In another, boracite is decomposed with the equivalent amount of sodium bisulphate (a by-product of the intric acid manufacturing process), the boric acid crystallised out, and mother-liquor worked up for the sodium sulphate it contains. The American deposits of colemanite are converted manily into boras.4

Orthoboric acid crystallises from water in white, six-sided lamine which have a pearly lustre and are unctuous to the touch—The crystals are triclinic (a b c 1.7329 1 0 9228, a 92° 30′,  $\beta$  101° 25′,  $\gamma$  89° 19′). According to Carnelley, orthobonic acid melts at 184° to 186′ "The density at 15° is given by Stolba  $^7$  as 1 434 . Ditte  $^8$  gives the following values

The specific heat is 0 3535 9

Orthoboric acid is sparingly soluble in cold water, but the solubility rapidly increases with rise of temperature. The percentages of orthoboric acid in its saturated aqueous solutions at various temperatures are as follows - 10

According to Herz and Knoch, 11 a saturated solution of orthoboric acid contains 0 620 gram-molecules of acid per little at 13°, 0 7915 at 20°, and 0 8999 at 25°. The cryohydric point is -0.76°, at which temperature the saturated solution contains 2 27 per cent of orthoboric acid. The boiling point of a

Nasim, Alti R. Acad. Lines, 1908, [v], 17, ii. 43; cf. D'Achaide, ibid., 1908, [v.],
 ii. 238, and the early papers of Bolley, Annalin, 1848, 68, 122; Waitington, Chemitagette, 1854, 7, 419; Wohler and Deville, Annalin, 1858, 105, 69; Popp, Annalin, Suppl., 1872, 8, 1, Diculatart, Ann. Chim. Phys., 1877, [v.], 12, 318; Compt. rend., 1885,

Suppl., 1872, 8, 1, Deculatat, Ann. Chem Phys., 1877, [v.], 12, 318, Compl. rend., 1885, 100, 1017, 1210.

Nasun, Anderlun, and Levi, Alti. R. Accad. Linect., 1905, [v.], 14, in. 70.

Nasun, Anderlun, and Levi, Alti. R. Accad. Linect., 1905, [v.], 14, in. 70.

Nasun, Anderlun, and Levi, Alti. R. Accad. Linect., 1905, [v.], 14, in. 70.

Nasun, Anderlun, and Levi, Alti. R. Accad. Linect., 1905, [v.], 14, in. 70.

Nasun, Anderlun, and Levi, Alti. R. Accad. Linect., 1905, [v.], 189, 189, 189, 180.

Narquardt and Schulz, Zeitsch. anger. Chem., 1895, 8, 385, and Thomps, A. Dictionary of Applied Chemistry (Longmans & Co., 1912-13), 2nd ed., vol. 1, p. 501.

Masshofer, Zeitsch. Kipst. Min., 1884, 9, 77, cf. Miller, Poogs. Annalen, 1831, 23, 1818, 1819, 1818, 1819, 1818, 1819, 1

saturated solution of orthoboric acid is 103.12°.1 The heat of solution of orthoboric acid in water is -5.395 Cals.<sup>2</sup> A saturated solution of orthoboric acid has a density of 1 014 at 8°, and 1 0218 at 15° (Stolba).

The solubility of orthoboric acid in water is increased by the presence of potassium or rubidium chloride, but diminished by the presence of hydrogen, lithium, or sodium chlorides 3 The and is very slightly soluble in other, more soluble in alcohols and essential oils.4 One hundred grams of glycerol dissolve the following amounts of orthoboric acid - 5

Temp. 
$$^{\circ}$$
 C. .  $0^{\circ}$  20 $^{\circ}$  40 $^{\circ}$  60 $^{\circ}$  80 $^{\circ}$  100 $^{\circ}$  Grams  $H_3BO_3$  20 28 38 50 61 73

The partition-coefficient of bonic and between water and ether is 31.2 at 16°; between isobutyl alcohol and water 2.74, and between amyl alcohol and water 3.37 at 15°, 3 34 at 25°, and 3.31 at 35°.6

Orthoboric acid is volatile in steam. The vapour given off from a saturated boiling aqueous solution of the acid contains 0 039 per cent. of acid.7 The acid may also be volatilised from its alcoholic solutions 8

The salts of borne and are called borates. Very few orthoborates are known, but numerous meta- and pyro-borates have been prepared. Further, salts of such hypothetical acids as II<sub>4</sub>B<sub>2</sub>O<sub>5</sub>, II<sub>6</sub>B<sub>8</sub>O<sub>15</sub>, etc., are known. Anhydrous borates may be prepared by fusing boron sesqui-oxide with metallic oxides. When an excess of boron sesqui-oxide is present, it is found that in some cases a homogeneous liquid mass is obtained which solidifies to a homogeneous glass; in other cases it separates on cooling into conjugate liquid phases; while in others it is not possible to obtain a homogeneous liquid melt, the mass separating into two non-miscible phases of metallic borate and boron sesqui-oxide respectively.9

The borates, with the exception of the alkali borates, are practically insoluble in water. The most important borate is sodium pyroborate or borax. The important borates are described under the headings of the various metals.10

Although the metallic orthoborates cannot be obtained by precipitation,

<sup>1</sup> Nasını and Ageno, loc cit, and Attı R. Accad. Lincer, 1912, [v ], 21, n 125. <sup>2</sup> Thomsen, Thermochemische Untersuchungen (Leipzig, 1883), vol in. p. 196; cf. Ditte, loc. cit.

Herz, Zeitsch. anorg. Chem., 1910, 66, 358.

<sup>\*\*</sup> Herz, Zeitsch. anorg. Chem., 1910, 66, 358.

\*\* Rose, Pogg. Annalen, 1850, 80, 262

\*\* Hooper, Pharm. J., 1882, [111], 13, 258; see also Herz and Knoch, loc. cit.

\*\* Abegg. Fox, and Herz, Zeitsch. anorg. Chem., 1903, 35, 129, Auchbach, ibid., 1904,
34, 353; B. Muller and Abegg, Zeitsch. physikal. Chem., 1907, 57, 513.

\*\* Nasını and Ageno, loc. cit., Skirrow, Zeitsch. physikal Chem., 1901, 30, 84.

\*\* See Futh and Myers, Trans Chem. Soc., 1914, 105, 2887.

\*\* Geurtler, Zeitsch. anorg. Chem., 1904, 40, 225, 268, 337; cf. Holt, Proc. Roy. Soc., 1902, 74, 285; Burgess and Holt, Proc. Chem. Soc., 1903, 19, 221; Proc. Roy. Soc., 1904, 74, 285.

\*\*For speculations on the constitution of borates, see Zulkowski, Chem. Zeitsch. 1900, [v.], 7, 1041; Ditte, Compt. rend., 1873, 77, 788; Le Chatelier, ibid., 1831, 113, 1034.

\*\*Mumerous borates are described in the following early papers, but the compositions of many of them are doubtful: Berzelius, Pogg. Annalen, 1824, 2, 113; 1827, 9, 438; 1834, 33, 98; 1835, 34, 561; H. Rose, ibid., 1827, 9, 176; 1830, 19, 153; 1852, 86, 561; 1852, 47, 1, 470, 587, 1853, 88, 299, 482; 1853, 89, 473; 1854, 97, 452; Wohler, ibid., 1833, 28, 28, 285; Rammelsberg, ibid., 1840, 49, 445; Ebelmen, Ann. Chim. Phys., 1851, [in.], 33, 34; Laurent, ibid., 1838, [ii.], 67, 215; Bolley, Annalen, 1848, 68, 122; Horepath, Phil. Mag., 1849, [iii.], 34, 375; Tissier, Compt. rend., 1854, 39, 192; 1857, 45, 411; C. L. Bloxam, Quart J. Chem. Soc., 1859, 12, 177; 1861, 14, 143; Ditte, Compt. rend., 1873, 77, 783, 892; Ann. Chim. Phys., 1853, [v.], 30, 248.

and have only in a few cases been prepared in other ways, the tri-alkyl esters of boric acid, B(OR), where R is an alkyl group Call min, are readily prepared by the interaction of boron trichloride and alcohols, eg:—

$$BCl_3 + 3C_2H_5 OH = B(OC_2H_5)_3 + 3HCl.$$

They are colourless liquids of low boiling-point and normal vapour density, and are readily hydrolysed by water! They combine with the metallic derivatives of the alcohols, forming crystalline compounds in which boron is presumably a pentad. The following are known ---

$$\begin{array}{lll} \operatorname{LiB}(\mathrm{OMe})_1 & \operatorname{KB}(\mathrm{OMe})_1 & \operatorname{NaB}(\mathrm{OEt})_1 & \operatorname{TiB}(\mathrm{OEt})_1 \\ \operatorname{NaB}(\mathrm{OMe})_1 & \operatorname{Ca}[\operatorname{B}(\mathrm{OMe})_4]_2 & \operatorname{KB}(\mathrm{OEt})_1 & \operatorname{NaB}(\mathrm{OP}_1^{\alpha})_1 \end{array}$$

In aqueous solution, orthoboric acid is a very weak acid. It turns litmus a wine-red, turmeric a reddish-brown, and has no effect on methylorange (p 44). The aqueous solution is a very poor conductor of electricity, so that the acid is ionised only to a very slight extent. The molecular weight of the acid in solution is, in fact, that corresponding to the molecular formula  $H_{a}BO_{b}^{-3}$  That there is only one molecular species present to any appreciable extent is shown by the fact that the partition-coefficients previously given (p 32) are independent of the actual concentration of boric acid in the aqueous phase. In solution, orthoboric acid behaves as a monobasic acid, The ionisation of the acid follows Ostwald's dilution law for binary electrolytes (Vol. 1. p 221), and the dissociation must therefore be represented as  $H_4BO_{q+}$  - $H_+H_2BO_4$ . The ionic mobility of the amon at 18 is 28.4 The admits constant (k) of orthoboric acid varies with the temperature as follows ---

Hence borne acid is weaker than either carbonic or hydrosulphuric acid. Owing, however, to the superior volatility of these acids, a concentrated, boiling solution of boric acid can decompose certain metallic carbonates and sulphides "

Both metaboric acid and pyroboric acid (orthoboric acid dehydrated till it corresponds to the formula, H<sub>2</sub>B<sub>1</sub>O<sub>7</sub>), when dissolved in water, are converted into orthoboric acid, since their aqueous solutions are identical with solutions

Ebelmen, Ann. Chim. Phus., 1816, [m], 16, 129; Ebelmen and Bouqet, ibid., 1846, [m.], 17, 54; Bowman, Phil May, 1816, [m], 29, 546, Rose, Pogy Annala, 1856, 98, 245; Schult, Annala, Suppl., 1867, 5, 151, Innala, 1877, 189, 162, Counclet, Ber., 1876, 9, 485; 1877, 10, 1655; 1878, 11, 106. J. prakt. Chem., 1878, [n], 18, 371. Ghirs, Gazzetta, 1193, 23, 1. 452, 11. 8, Copaux, Compt. rend., 1898, 127, 719, Wohl and Neuberg, Ber., 1899, 32, 3483. Fenton and Miss Gostling, Trans. Chem. Soc., 1668, 73, 554; Cohn, Pharm. Zentr. h., 1911, 52, 479
 Copaux, loc cit., Cambi, Att. R. Accad. Lincei, 1914, [v.], 23, 1. 244
 Kahlenberg and Schreiner, Zeits-h. physikal Chem., 1806, 20, 547; Holt, Mem. Man. Phil. Soc., 1911, 55, No. 10; Nasini and Ageno, loc. cit.
 Abegg and Cox, Zeitsch. physikal Chem., 1903, 46, 1, Walker and Cormack, Trans. Chem. Soc., 1900, 77, 5
 Lundén, J. Chim. phys., 1907, 5, 580; cf. Walker and Cormack, Trans. Chem. Soc., 1900, 77, 5;
 Hantsch and Barth. Ber., 1902, 35, 216, concentration in gram molecules per litre.

per litre.

Tissier, Compt. rend., 1854, 39, 192, 1857, 45, 411, Popp, Annulen Suppl., 1872, 8, 1. VOL. IV.

of orthoboric acid of the same boron content. Moreover, aqueous solutions of alkali meta- and pyro-borates are in every respect identical with aqueous solutions prepared from the requisite quantities of orthoboric acid and alkali \_ hydroxide. In conformity with the view already outlined that boric acid in solution is a monobasic acid, it reacts in aqueous solution with one equivalent of alkali. This has been shown both by freezing-point measurements and by thermochemical considerations. Accordingly, the ions in alkali borate solutions are essentially M and H<sub>2</sub>BO<sub>3</sub> or BO<sub>2</sub>, and such solutions, by double decomposition with metallic salts, precipitate sparingly soluble metaborates. There must, however, be an approxiable concentration of hydroxyl ions in an alkali borate solution, since, bone and being exceedingly weak and the alkali hydroxides very strong, the salts produced by their interaction must be perceptibly hydrolysed (Vol I. p. 221). At ordinary temperatures, in fact, a decinormal solution of borax is hydrolysed to the extent of about 0.5 per cent.3 Accordingly, in sufficiently dilute solutions, precipitated metallic metaborates are contaminated with co-precipitated metallic hydroxides (or oxides) if the hydroxides are very sparingly soluble substances 1. Alkali borate solutions also contain complex polyborate ions, the precise nature of which is not known.5

Boric acid and its most important salt, borax, receive many practical applications. As a mild antisoptic, the acid is largely used as a food preservative. It is also used in the preparation of candle wicks. Boric acid and borax are employed in the preparation of enamels, pottery glazes, hat varnishes, paint direis, borosilicate glass, certain kinds of optical glass, cosmetics, tooth powders, soaps, parchiment paper, glazed paper and caids, safe linings, and fireproof textile fabrics, they are also used by tanners for dressing leather, by coppoismiths and jewellers as fluxes in brazing and soldering operations, by laundresses, etc.

Complex Boric Acids.—Complex acids containing boron are formed by the action of boric acid upon numerous substances. The best defined inorganic compounds of this nature are the two borotungstic acids, [6H<sub>2</sub>O.B<sub>2</sub>O<sub>3</sub> 28WO<sub>3</sub>].56H<sub>2</sub>O and [5H<sub>2</sub>O.B<sub>2</sub>O<sub>3</sub> 24WO<sub>3</sub>].61H<sub>2</sub>O, prepared by Copaux.6

The interaction of boric acid with numerous organic compounds has been studied by Magnanin, who found that in aqueous or alcoholic solution, boric acid interacts with all hydroxycarbovylic acids in which at least one hydroxyl group is present in the a-position to a carbonyl group, the complex acids formed (which he did not isolate) being stronger than the organic acids from which they are produced. The other organic acids do not interact

Noyes and Whitney, Zeitsch. physikal. Chem., 1894, 15, 694.
 J. Thomsen, Thermochemische Untersunhungen (Leipzig. 1882), vol. 1. p. 206; cf. Iandén, J. Chim. phys., 1907, 5, 599.
 Shields, Zeitsch. physikal. Chem., 1893, 12, 167.
 Abegg and Cox, loc. ett.; cf. the interaction of borax and silver nitrate, p. 44.
 Auerbach, loc. ett.; P. Muiler and Abegg, loc. ett.
 Copaux, Compt. rend., 1908, 147, 973, Ann. Chim. Thys., 1909, [viii.], 17, 217.
 See Vol. VII. of this series. For evidence in lavour of the existence of boro molybdates, boroarsenates, etc., see Mauro, Ber., 1881, 14, 1379; Bull. Soc chim., 1880, ii.], 33, 564;
 Rosenheim and Bertheim, Zeitsch. anorg. Chem., 1903, 37, 353.
 Magnanini, Atti. R. Accad. Lincer, 1890, [iv.], 6, 1. 260, 411, 457; Gazzetta, 1890, 280, 428, 441, 448, 453; 1891, 21, ii. 184, 215; 1911, 41, ii. 425, 1914, 44, ii. 396; Zeitsch. physikal. Chem., 1890, 6, 58; 1892, 9, 230; 1898, 11, 281.

with boric acid in solution. A crystalline potassium hydrogen borooxalate 2K(BO)C2O4 3H2O,1 can nevertheless be readily prepared by heating potassium metaborate with oxalic acid solution, or by heating boric acid

with potassium hydrogen oxalate

The interaction of boric acid with alcohols has been studied by Magnanin, P. Muller and Abegg, and others.<sup>2</sup> It has been shown by Muller and Abegg that complex acids are formed (by condensation of the alcohols with boric acid) to an appreciable extent when boric acid is dissolved in various hand alcohols, but that the extent to which complex acid formation occurs, steadily diminishes as the alcoholic solvent is replaced by water. Complex acid formation between borie acid and both mannitol and glycerol, two polyhydric alcohols, is of considerable interest, both theoretically and practically. In each case the complex acid formed is much stronger than boric acid, and whereas mannitol is optically mactive, mannitoboric acid is decidedly dextro totatory. The extent to which these complex acids are formed in solution increases with the concentration of either the boric acid or the polyhydric alcohol, and diminishes with the dilution, in consequence of which the molecular conductivity of mannitoboric acid decreases with the dilution. Magnanini concluded from his experiments that mannitoboric acid was produced by the condensation of three molecules of bone and with one of mammitol, but the subsequent isolation of the complex acid has shown that the condensation proceeds as expressed by the equation -

$$H_1BO_3 + C_6H_{14}O_6$$
.  $C_6H_{15}O_5B + H_2O_6$ 

The preparation of mannitoboric acid is carried out by dissolving 12 grams of mammitol and 8.2 grams of bone acid in 95 cubic centimetres of absolute alcohol, and setting the solution aside for a few days in a cold place. The first few crystals formed are discarded, those subsequently deposited are pure manufoboric acid, CaH15OaB, and melt at 89.5". The molecular weight of the acid in acctone solution is rather greater than that required for the preceding formula. The acid is largely decomposed by water, but its ammonium, silver, calcium, and barium salts are more stable <sup>3</sup>

Perboric Acid and Perborates.-Perboric acid is unknown in the pure state, but it is possible that ethereal solutions have been prepared. A number of metallic perborates, however, are known. They were discovered by Etard, and first prepared in a pure state by Melikoff and Pissarjewsky.

The best known perborates are derivatives of a perbone acid of the composition HBO. They are readily hydrolysed in aqueous solution, boric acid, or rather its salts, and hydrogen perovide being produced. Accordingly, the perboric acid is considered to have the constitution O BOOH :-

O BO OH +  $2H_2O = B(OH)_3 + HO OH$ .

E. A. Werner, Trans Chem. Soc., 1901, 85, 1449.
 P. Muller and Abegg, Zeitsch physikal. Chem., 1907, 57, 513; Ageno and Valla, Gazzetta, 1913, 43, ii. 163, Dahr, Zeitah. ano. g. Chem., 1914, 86, 196; Boeseken and others, Rec. trav. chem. 1911, 30, 392; 1915, 34, 96, 272, Proc. K. Akad. Wetensch. Amsterdam, 1912, 15, 216, Irvine and Miss Stycle, Trans. Chem. Soc., 1915, 107, 1221.
 Fox and Gauge, Trans. Chem. Soc., 1911, 99, 1075.
 Pissarjewsky, Zeitsch. physikal. Chem., 1903, 43, 170.
 Etard, Compt. rend., 1880, 91, 981.
 Melikoff and Pissarjewsky, Ber., 1895, 31, 678, 953.

This constitution, however, has been disputed on the ground that hydrated podium or potassium perborate does not give off hydrogen peroxide when heated to 50'-60° under diminished pressure or in a current of air free from

carbon dioxide; and the alternative constitution HO.B. has been pro-

posed by Bosshard and Zwicky  $^1$  . The following argument, based upon the preparation and properties of  $\rm KBO_p$  is also given by the same chemists. The compound KBO, or polassium hyperborate, crystallises with 1H2O and is fairly stable. When dried over phosphoric anhydride in vacuo, 4KBO, H2O is produced, so that the original compound cannot be formulated as KBO, H<sub>2</sub>O<sub>2</sub>. It is rather to be considered as derived from potassium hyperoxide KO OII, and perboric acid. Hence, two constitutions are possible, viz KOO.OB.O

and KO ,  $0.8 \stackrel{O}{\underset{\sim}{\bigcirc}}$  , and it is considered that the latter is much more probable

than the former owing to the stability of the salt

Sodium perborate, NaBO, 111,0, may be prepared by adding to a saturated solution of borax an equivalent quantity of sodium hydroxide and twice the amount of hydrogen perovide theoretically necessary, after some time the perborate commences to crystallise out - 2

$${\rm Na_2B_4O_7 + 2NaOH + 4H_2O_2 - 4NaBO_4 + 5H_2O}.$$

In preparing the salt on a large scale the necessary ingredients are mixed in small batches, which are then united and slowly cooled with stirring Wooden vats and strices are used, and metal pipes, etc, well timed to minimise decomposition of the hydrogen peroxide. The temperature after mixing is 25°, and the concentrations are chosen so that the perborate separates out slowly. Large and fauly stable crystals are deposited, which are centrifuged free from mother liquor and dried in a stream of pure air at a temperature not exceeding 50° C 3

Sodium perborate is only formed in minute quantities at the anode when aqueous sodium orthoborate, with or without excess of alkali, is electrolysed.4

Sodium perborate, NaBO, 4H,O, crystallises in large, transparent, monoclinic prisms, and is stable in air free from carbon dioxide 5. The monohydrate may be obtained by careful drying, and this may be completely dehydrated in vacuo over phosphoric anhydride The solubility of the salt in water, in grams per litre of solution, is as follows .--6

Тешр. ° С	15°	$21^{\circ}$	26	35,
Grams of NaBO, 4HaO	25.5	26.9	28.5	37.8

Bosshard and Zwicky, Zeitsch angew. Chem., 1912, 25, 993
 Melikoff and Presaijewsky, loc. cit.; Tanatri, Zeitsch physikal Chem., 1898, 26, 132;
 1899, 29, 162; Jaubert, Compt. rend., 1904, 139, 796; Bruhat and Dubois, ibid., 1905, 140,
 506; Jaubert and Lion, Rev gen. chem pure appl., 1905, [vn.], 8, 163, Christensen,
 Danske Kidensk Seisk Fonkandt, 1904, No. 6.
 Fuhrmann, Chem. Zeit., 1911, 35, 1022.
 See Tanatar, Zeitsch. physikal. Chem., 1898, 26, 132; Zeitsch. anory. Chem., 1901, 26,
 Sta, Constam and Bennett, ibid., 1900, 25, 265., 1901, 26, 461, Bruhat and Dubois, Compt. rend., 1905, 140, 506; Presaijewsky, Zeitsch anory. Chem., 1902, 32, 311; Poulene, Fr. Pat., 411, 268; Beltzer, Moniteur Scient., 1911, p. 10, Polack, Trans. Faraday Soc., 1915, 10, 177.

<sup>1915, 10, 177.</sup>Jaubert, Compt. rend., 1904, 139, 796.
Jaubert and Lion, loc. cit.

The solubility is augmented by the addition of borie, tartanic, or citric acid, glycerol, or ammonium and magnesium sulphates in small amount 1 The heat of solution in water is  $-11^\circ56$  Cals at 16", in seminormal sulphuric acid it is  $-8\,95$  Cals at 17° C.2

In aqueous solution at 0° C the hydrolysis of sodium perborate is very slight, and from conductivity measurements it follows that the molecular formula of the salt is  $NaBO_3$ . With rise of temperature, hydrolysis proceeds further, and at 25° it is very considerable. The aqueous solution has an alkalme reaction. According to Pissarjewsky, the hydrolysis proceeds in two stages, (i) and (ii), the third reaction (iii) then taking place between products formed in (1) and (11.)

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(1) NaBO_3 + 2H_3O_3 = NaO_3H + H_3BO_3
(ii.) VaOOH + H_0O = -NaOH + H_0O,
(iii.) 3\text{NaOH} + \text{H}_4\text{BO}_4 = -\text{Na}_4\text{BO}_4 + 3\text{H}_2\text{O}_4
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When heated above 40°, sodium perborate solution commences to evolve ovygen. The solution is an energetic oxidiser, converting ferrous salts to ferric, chromic salts to chromic acid, manganous salts to manganese dioxide, liberating chlorine from hydrochloric acid, iodine from potassium iodide, etc. 5

Sodium perborate is used commercially under the name of "perborin" for bleaching purposes, since it may be incorporated into soaps and washing powders "Perborm M" contains soap, alkali, and sodium perborate; persil" contains soap, alkali carbonate and silicate, and sodium perborate; "ozonite" is similar to "persil', and "clarax" contains borax, sodium phosphate, and sodium perborate 6

A sodium perborate of the composition NaB2O1 5H2O is said by Jaubert to be obtained when an intimate mixture of 248 grams of boric acid and 78 grams of sodium peroxide is slowly added to ! littles of cold water. It cannot be recrystallised from water

Sodium hyperborate, NaBO, has not been prepared in a pure state 8 Potassium perborate. The following salts have been described (i.) KB<sub>3</sub>O<sub>5</sub> 2H<sub>2</sub>O<sub>5</sub> (in.) 2KBO<sub>4</sub> KBO<sub>4</sub> 5H<sub>2</sub>O or 3KBO<sub>4</sub> H<sub>2</sub>O<sub>5</sub> 4H<sub>2</sub>O<sub>5</sub>(in.) 2KBO<sub>3</sub> H<sub>2</sub>O<sub>5</sub> and (iv.) 2KBO3.H2O2.11

Rubidium perborate, RbBO, II2O, and cæsium perborate, CsBO, H2O, are also known.12

Ammonium perborate, NH<sub>1</sub>BO<sub>1</sub> 3H<sub>2</sub>O<sub>2</sub> is obtained by dissolving boric

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1 Bruhat and Dubors, loc cit. , Jaubert and Lion, loc cit
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Bruhat and Dubois, loc cit., Jaubert and Lion, loc cit
 Tanatai, Zeitsh phusikal Chem., 1898, 26, 132
 Constain and Bennett, Zeitsh among Chem., 1900, 25, 265
 Prisarjewsky, Zeitsh phusikal, Chem., 1903, 43, 170, cf. Schenck, Vorlander, and Dux, Zeitsh angew Chem., 1911, 27, 291.
 For further ractions, see Jaubert, Compterend., 1904, 139, 796, Bruhat and Dubois, Compterend., 1905, 140, 506, Christensen, Danske Vidensk Sidsk Ferhandl., 1904, No. 6.
 Matthews, J. Ind. Eng. Chem., 1911, 3, 191
 For the patent literature dealing with the manufacture of alkali perhorates, see D.R.P., Nos. 193,559, 204,279, 237,096, 237,608; D.R.P. Amn., 19,641, Fr. Pat., Nos. 384,967, 411,258, 425,958, U.S. Pat., Nos. 996,778, 999,497; Eng. Pat., 1626 (1911), cf. Bosshard and Zwicky, Zeitsch angew. Chem., 1912, 25, 938

 <sup>325, 938</sup> Jaubert, Compt. rend., 1994. 139, 796
 Melikoff and Pissarjewsky, Br. 1898, 31, 678, 953.
 Bruhat and Dubois, Compt. rend., 1995, 140, 596.
 Christensen, Danske Vidensk. Selsk. Forhandl., 1994, No. 6.
 Yon Girzewald and Wolokilin, Ber., 1999, 42, 865.

<sup>12</sup> Christensen, loc. cit.

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acid in 2.5 per cent. hydrogen peroxide, adding ammonia, and precipitating with alcohol. When dued for twenty-four hours over sulphuric acid, the salt 2NH<sub>4</sub>BO<sub>3</sub>·H<sub>2</sub>O is obtained, and this may be dehydrated over phosphoric anhydride.<sup>1</sup>

The salt NH, BO<sub>3</sub>, NH<sub>1</sub> BO<sub>1</sub> 2H<sub>2</sub>O has been described by Petrenko <sup>2</sup>

Barium perborate, Ba(BO<sub>3</sub>)<sub>2</sub> 7H<sub>2</sub>O, <sup>3</sup> and uranyl perborate, UO<sub>2</sub>BO<sub>3</sub>, <sup>4</sup>

are also known.

Boron sesquisulphide (boron trisulphide), B<sub>2</sub>S<sub>3</sub>, is produced when boron is heated to bright redness in sulphur vapour,<sup>5</sup> or when boron is heated to redness in a current of hydrogen sulphide <sup>6</sup> It may also be prepared by heating to redness in a current of carbon disulphide vapour an intimate mixture of boron sesqui-oxide and carbon <sup>7</sup> Other reactions which lead to the formation of the sulphide are the action of sulphir upon boron tri-odide and the action of carbon disulphide and various other sulphides on boron (Moissan). The pure sesquisulphide is best prepared by heating pure boron in a stream of hydrogen sulphide largely diluted with hydrogen, and a cheap method for its preparation consists in heating commercial iron boride or manganese boride to 300°-100° in hydrogen sulphide and washing away admixed sulphir from the product with carbon disulphide. Only part of the boron in these borides can be removed as the sulphide

The properties of boron sesquisulphide have been studied mainly by Moissan. It forms fine, white needles, of density 155, which melt at 310° and volatilise without decomposition when heated in a current of hydrogen. It is unacted upon at a red heat by hydrogen, introgen, phosphorus, carbon, and silicon—It inflames in chlorine in the cold, in bromme vapour when warmed, and in oxygen when heated to dull redness, in each case both the boron and sulphur are converted into chloride, bromide, or oxide. At a dull red heat, boron sesquisulphide is violently decomposed by potassium, sodium, magnesium, and aluminium, but non, zinc, copper, mercury, and silver are without action upon it—It is decomposed by carbon dioxide above 300° as follows:—11

$$B_2S_3 + 3CO_2 = B_2O_3 + 3CO + 3S_4$$

It is violently decomposed by water at ordinary temperatures, boric acid and hydrogen sulphide being produced and much heat evolved --12

$$[B_2S_3] + 6H_2O + aq. = 2H_3BO_3aq. + 3(H_2S) + 57.8 \text{ Cals}$$

Boron sesquisulphide is slightly soluble in phosphorus trichloride, from which it crystallises in needles. It forms the addition-compounds B<sub>2</sub>S<sub>4</sub>.BCl<sub>2</sub>

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1 Melikoff and Pissarjewsky, Ner., 1898, 31, 678,953; Constam and Bennett, Zeitsch. anorg. Chem., 1900, 25, 265; Tanatar, Zeitsch. physikal. Chem., 1898, 26, 132; Bruhat and Dubois, Compt. vend, 1905, 140, 506

2 Petrenko, J. Russ. Phys. Chem. Soc., 1902, 34, 37.

3 Melikoff and Pissarjewsky, loc. cit.

4 Bruhat and Dubois, Compt. rend, 1905, 140, 506.

5 Berzelits, Pog. Annalen, 1824, 2, 145.

5 Wohler and Deville, Ann. Chim. Phys., 1858, [iii.], 52, 90.

7 Frómy, Ann. Chim. Phys., 1853, [iii.], 38, 312.

8 H. Gautier, Ann. Chim. Phys., 1859, [vii.], 18, 363.

5 J. Hoffman, Zeitsch. angew. Chem., 1906, 19, 1362, 2133; Zeitsch. anorg. Chem., 1908, 59, 127.

1908, 59, 127.

1008, 59, 127.

1008 Moissan, Compt. rend., 1892, 115, 203.

11 Costeanu, Compt. rend., 1891, 112, 862.
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and B2S3.BBr3, and according to Moissan, combines with the chlorides of phosphorus, arsenic, and antimony. When dissolved in hot carbon disulphide, saturated with hydrogen sulphide, the solvent evaporated in racuo and the residue recrystallised from either carbon disulphide or benzene, white crystals of B.S. H.S or thiometaboric acid are obtained. This substance, which smells of hydrogen sulphide and is rapidly decomposed by water, has a molecular weight corresponding to the formula HaBaS, in benzene solutions; unlike boron sesquisulphide, it is extremely soluble in both benzene and carbon disulphide. Heated in a scaled tube, it begins to melt at 120° and forms a clear, transparent liquid at 140', when heated in air, it readily dissociates into boron sesquisulphide and hydrogen sulphide. Thiometaboric acid dissolves in liquid ammonia, when excess of solvent is removed at ordinary temperatures, yellow crystals of B2S3.6NH3 separate out.1

Boron pentasulphide, B.S. -When sulphur reacts upon boron tri-fodide without the intervention of a solvent, boron sesquisulphide is produced, but when the reaction occurs in carbon disulphide solution, crystals of boron pentasulphide are said to separate from the liquid 8. The substance dissociates into sesquisulphide and sulphin when heated in vacuo. It has not been obtained with less than 10 to 15 per cent of admixed iodine,

Boron selenide, B.Se, may be prepared by heating boron in selenium vapour or hydrogen selenide <sup>4</sup> It is also produced if iron or manganese boride be used instead of boron itself <sup>5</sup> The selenide is a yellow solid which does not appear to be volatile at a red heat. It is decomposed by water in a similar manner to the sulplide

# BORON AND THE NITROGEN GROUP.

Boron nitride, BN -Boron unites directly with nitrogen at a red heat to form a nitride (Wohler and Deville, Moissan) Boron mitride was discovered by Balmann' in 1842 by fusing boron sesqui-oxide with potassium cyanide, but its true nature was determined and the compound prepared in a nearly pure state by Wohler in 1850,7 His method consisted in heating an intimate mixture of anhydrous borax (1 pt.) and ammonium chloride (2 pts ) to bright redness in a platinum crucible, boiling the powdered product with very dilute hydrochloric acid as long as boric acid could be extracted, washing and drying the residue. Another method of preparation consists in heating a mixture of boion sesqui-oxide (4 pts.) and carbon (1 pt ) to whiteness in a current of introgen.5 The nitride is produced when boton is heated to redness in ammonia, and may be more readily prepared by heating either boron sesqui-oxide or anhydrous borax in the same gas 9 Numerous other methods have been proposed, as, for example, heating boron sesqui-oxide or borax with potassium ferrocyanide, potassium ... cyanide, mercuric cyanide, or urea 10

<sup>1</sup> Stock and Blv, Ber., 1901, 24, 3039, 1903, 36, 819.
2 Stock and Poppenberg, Ber., 1901, 34, 399.
3 Moissan, Compt. rend., 1892, 115, 271.
4 Sabatier, Compt. rend., 1891, 112, 1000.
5 J. Hoffmann. Chem. Zett., 1911, 35, 713.
6 Balmann, Phit. Man., 1812, [nn.], 21, 170, 1843, [nn.]. 22, 467, 1844, [ini.], 24, 191.
7 Wohler, Annalen, 1850, \$4, 70
8 Wohler and Deville, Ann. Chim. Phis., 1858, [ini.], 52, 63
9 See Ehrich and Graetz, D.R.P., 282,701 (1913); Podsvus, D.R.P., 282,748 (1918).
10 Darmstadt, Annalen, 1869, 151, 255, Martius, while, 1859, 109, 80; Rose, Pogg., Annalen, 1850, 80, 265; Gustavson, Ber., 1870, 3, 426.

As a convenient method of preparation, Moeser and Edmann 1 recommend passing ammonia over a strongly heated and previously fused mixture of boron sesqui-oxide and calcium phosphate. The product, when washed with dilute hydrochloric acid, is nearly pure boron intride. To obtain the nitride in a state of purity, boron tribromide should be slowly dropped into liquid ammonia, the excess of solvent removed by evaporation, and the solid residue gradually heated to 750° 2 The made of boron unitally formed (p. 24) becomes converted into nitride

$$B_2(NH)_1 - 2BN + NH_3$$
.

The method given by Wohler and Deville for the preparation of boson nitride has been studied by Stahler and Elbert, with respect to its utilisation for the fixation of atmospheric nitrogen. When a mixture of boron sesquioxide and carbon is heated in nitrogen, the yield of boron introgen varies with the temperature and the pressure of the introgen At a pressure of one atmosphere, the best yield, 26 to 28 per cent. of intride, is obtained at 1500° to 1700°; but at a pressure of 70 kilos per square centimetre, a yield of more than 85 per cent. is obtained at 1600°. When, however, boron sesqui-oxide is replaced by borocalcite, CaB<sub>1</sub>O<sub>7</sub>, a nearly theoretical yield of boron nitride, according to the equation

$$CaB_1O_7 + 8C + 3N_2 = -4BN + CaCN_2 + 7CO_5$$

is obtained, even at one atmosphere pressure, by heating first to 1850° and subsequently lowering the temperature to 1400°; increasing the pressure of the nitrogen has practically no effect on the yield.

Boron nitride is a light, white, amorphous solid, soft, like tale, to the touch It can be compressed, wet or dry, into blocks of considerable nigidity. It is infusible at the melting-point of tungsten, but above 1500 it commences to dissociate in vacuo into boron and nitrogen 4. At 1220', however, the dissociation pressure does not exceed 9.1 min. 5. At high temperatures it is the best insulator known 6 Heated in a flame it exhibits a greenish-white fluorescence

Boron nitride is a very stable compound. It is very slowly decomposed by boiling water, aqueous potash, hydrochloric and nitric acids, but decomposes more readily when heated in steam or at 200° in a scaled tube with water, hydrochloric or sulphure acid -

$$BN + 3H_3O - H_3BO_3 + NH_{30}$$

Heated with fused potassium hydroxide, ammonia and potassium borate are produced, with fused potassium carbonate the products are potassium cyanate, cyanide, and borate. It is little affected by heating with oxygen, iodine, hydrogen, carbon dioxide, or carbon disulphide In an alcohol flame fed with oxygen it burns to boron sesqui oxide, and at high temperatures it is-decomposed by chlorine. It is slowly dissolved by hydrofluoric acid, ammonium borofluoride being produced. The oxides of copper, cadmium,

Mocser and Eidmann, Ber., 1902, 35, 535.
 Stock and Blix, Ber., 1901, 34, 3039; Stock and Holle, ibid., 1908, 41, 2095.
 Stahler and Eibert, Ber., 1913, 46, 2060.
 See Eng Pat., 25, 978 (1906).
 Slade and Higson, Rep Brit Assoc, 1913, p. 451.
 Weintraub, J. Ind. Eng. Chem., 1913, 5, 166.

mercury, arsenic, antimony, and bismuth are reduced when heated with boron mitride, the products being metal, metallic borate, and mitrous oxide; sulphates are reduced to sulphides, but the oxides of vine and iron are not reduced

100

Boron intride prepared by heating boron imide at 125° to 130° for a long time is, according to Stock and Blix, much more reactive than the nitride prepared at high temperatures, and they suggest that the latter is a polymer of the former

Boron amide, B(NH<sub>2</sub>), —The preparation of this compound has been already described (p. 23).—It has not yet been prepared in a state of purity; the accompanying ammonium salt can be removed partly, but not entirely, by washing with liquid ammonia

Boron imide, B2(NII), is best prepared by heating the compound B,S, 6NH, at 115° to 120° m a rapid stream of ammonia for some days -

$$B_2S_4 6NH_1 - B_2(NH)_1 + 3NH_1SH.$$

Other methods of formation have been already mentioned (pp. 23, 24)

Boron made is a light, white powder, insoluble in alcohol, ether, carbon disulphide, and liquid ammonia. It begins to decompose at 125" to \( \)30' with the evolution of ammonia, and at a slightly higher temperature is con pletely resolved into boron intride and ammonia. When boiled with water, boron imide is decomposed into ammonia and boric acid

Boron made is a feebly basic compound, and when added to liquid hydrogen chloride it forms a white hydrochloride, B2(NH), 3HCl, insoluble in organic media, and decomposed by heat or water

Boron phosphide, BP, is prepared by heating boron phospho-iodide to 450°-500' in a current of hydrogen. The indine can also be removed from the phospho-iodide by heating with mercury or silver. Another method of preparation consists in heating the compound BBr, PH, to 300', when hydrogen brounde is chimilated 4

Boron phosphide is a light, amorphous, maroon-coloured powder, insoluble in the usual morganic and organic solvents. It burns in chlorine in the cold and in bromine when warmed, at 200 at burns brilliantly in oxygen, and it also reacts with heated sulphur. It is not affected by iodine, introgen, phosphorus, or arseme even at a red heat. It is attacked by numerous metals at a red heat, and is easily oxidised by concentrated intric acid or fused alkalı nitrates

When heated in hydrogen at 1000°, a brown boron subphosphide, B<sub>5</sub>P<sub>9</sub> is produced which is much less reactive than the phosphide

Boron phospho-iodides .-- The compound BPI, is prepared by acting upon boron tri-iodide with phosphorus in carbon disulphide solution, and, in a carbon dioxide atmosphere, washing away the iodide of phosphorus simultaneously produced with a further quantity of carbon disulphide.<sup>5</sup> It is a dark red solid which melts in racuo at 190° to 200° and sublimes at higher temperatures. Chlorine, oxygen, and many metals attack it vigorously, and water immediately decomposes it, among the products being hydriodic,

Wohler and Deville, loc cit, Darmstadt, loc cit., Moeser and Eidmann, loc. cit.
 Stock and Blix, Brr, 1901, 34, 3039.
 Moissan, Compt. rend, 1891, 113, 726, 787.
 Besson, bid., 1891, 113, 78, 772.
 Moissan, Compt. rend., 1891, 113, 684.

♥ phosphorous, and boric acids. When heated to 160° in hydrogen it is reduced to the compound BPI, which sublines in vacuo, forming orange-yellow crystals, and closely resembles the other phospho-iodide in properties.

#### BORON AND THE CARBON GROUP.

Boron carbide, B.C —This compound was isolated in small quantities by Joly in 1883 1 from among the products of the interaction, in a carbon crucible, of boron sesqui-oxide and aluminium. Later it was prepared in quantity by Moissan, by heating to 3000' in the electric furnace a mixture of sugar carbon and amorphous boron contained in a carbon crucible. A cheaper method of preparation consists in employing boron sesqui-oxide and petroleum coke as starting materials, the former being fed through a hollow graphite electrode on to a layer of the latter contained in a graphite crucible which forms the other electrode 3 The product is purified from graphite by many treatments with boiling intric acid and potassium chlorate, followed by treatment with boiling sulphune acid

Boron carbide forms brilliant black crystals, of density 2.51, and melts at 2350.4 The carbide conducts electricity. It is extremely hard (harder than silicon carbide) and can be used in diamond polishing. It is not affected by mineral acids at the boiling temperature, sulphur at 500°, nitrogen, phosphorus, bromme, or nodme at a bright red heat. It is slowly attacked by oxygen at 1000°, and by chlorine at a somewhat lower temperature When fused with potassium hydroxide it is decomposed with the evolution of carbon monovide.

The so-called carbide 5 BC is a mixture of the preceding carbide and graphite (Tucker and Bliss).

Boron thiocyanate, B(CNS), is produced by shaking silver thiocyanate with a solution of boron tribionide in benzene, and separates from benzene in short, glistening, rhombic crystals. From other it crystallises in thin plates. The thiocyanate is rapidly hydrolysed by water, and its action upon aniline indicates that it possesses the structure of a thiocyanate rather than that of a throcar bundle.

Boron Alkyls,  $B(C_nH_{2n+1})_p$ —Although at present a hydride of the molecular formula BII, is unknown, the boron trialkyls have been known for many years. They are prepared by the action of zine alkyls on trialkyl esters of orthoboric acid. Boron trimethyl, B(CH<sub>3</sub>), is a gas at ordinary temperatures, boron triethyl, B(C<sub>2</sub>H<sub>3</sub>), is a volatile liquid. Both are readily combustible, burning with a green flame, and they combine readily with ammonia The vapour densities correspond with the above formulæ.

Boron Silicides, B.S. and B.S. These compounds are produced when , a mixture of crystalline silicon (5 pts ) and amorphous boron (1 pt ), packed in a refractory earthenware tube, is heated for 40 to 60 seconds by means of an electric current of 600 ampères at 45 volts. The outer portions of the

Joly, Compt. rend., 1883, 97, 456.
 Moissan, ibid., 1884, 118, 556.
 Tucker and Bliss, J. Amer. Chem. Soc., 1906, 28, 605; Tucker, D.R.P., 206,177;
 Tiede and Blimbrauer, Zeitch awry. Chem., 1914, 87, e129.
 Pring and Felding, Trans. Chem. Soc., 1909, 93, 1497.
 Muhlhauser, Zeitsch. anorg. Chem., 1893, 5, 92.
 Cocksedge, Trans. Chem. Soc., 1908, 93, 2177.
 Frankland and Duppa, Phil. Trans., 1862, 152, 167; Proc. Roy. Soc., 1376, 25, 165.

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product and those portions which were in proximity to the carbon electrodes are discarded, and the remainder treated with a cold mixture of nitric and hydrofluoric acids to eliminate free silicon. The residue is purified by heating for half an hour with commercial potassium hydroxide, which is just melted but not dehydrated After washing with water, dilute nitric acid, and boiling water, and drying at 130°, a mixture of the two silicides is obtained

The silicide B<sub>8</sub>Si is left when the mixture is heated with an excess of boiling mitric acid, the other compound being completely decomposed. It forms black, rhombic plates which are transparent and yellow or brown in very thin layers. The density is 2.52 lt easily scratches quartz and ruby, but is softer than boron carbide. Fluorine, chlorine, and bromine attack it in increasing order of difficulty, and it is only superficially oxidised by air or oxygen at a red heat. It is rapidly oxidised by fused anhydrous potassium hydroxide, less rapidly by fused alkali carbonates, and not at all by fused alkali intrates. Boiling concentrated sulphurio acid slowly oxidises it

The silicide B.Si is isolated from a mixture of the silicides, in which it is more abundant than the other, by heating with fused anhydrous potassium hydroxide It forms thick, opaque crystals of density 2 17. In chemical properties it resembles the other silicide, except in its behaviour towards fused potassium hydroxide and boiling nitric acid 1

# DETECTION AND ESTIMATION OF BORON.2

Boric acid, or a borate moistened with concentrated sulphuric acid, imparts a green colour to the flame The material to be tested should be introduced 2 centimetres above the top of the Bunsen burner, and should not be brought closer than 2 millimetres to the visible edge of the flame, in order that other substances may be prevented from tinging the flame green. coloration is best seen tangentially to the flame, and a hydrogen flame is preferable to the ordinary gas flame 3 The flame of a burning alcoholic solution of an alkyl borate is also coloured given (C. Geoffrey, 1732). To utilise this fact in qualitative analysis, the material under examination is mixed with concentrated sulphune acid and alcohol (preferably methyl alcohol), the alcohol ignited, and the mixture stirred 4 This test is not reliable in the presence of chloudes, phosphates, molybdates, barium, copper, thallium, and tellurium, and moreover, it fails to detect boron in many minerals Glycerol may be used instead of alcohol; in this case the substance is heated with a little sulphuric acid until excess of the latter has been driven off, the residue moistened with glycerol and a light applied.5 A more delicate and very characteristic test consists in heating the substance with calcium fluoride and concentrated sulphuric acid and leading the evolved gases into the Bunsen flame. A mere trace of boron trifluoride will tinge

<sup>&</sup>lt;sup>1</sup> Moissan and Stock, Compt rend , 1900, 131, 139 , Ann. Chim. Phys., 1900, [vii.].

<sup>20, 433.

&</sup>lt;sup>2</sup> A very full account of the analytical chainstry of boron, and copious references to the literature, will be found in Mellor, A Treatise on Quantitative Inorganic Analysis (Griffin & Co., 1913)

<sup>3</sup> Diculaint, Ann. Chim Thys., 1877, [v], 12, 318. See also Borntrager, Zeitsch. anal.

Chem., 1900, 39, 92

Dieulafast, loc cit.; Gilm, Rer., 1878, 11, 712; Lenher and Wells, vide infra.

Rev., 1877, 35, 204.

(L090) JS

(4985)A

(1585)11

(0685)\*Å (9685)\*N

(8049)

1500

ğ

Spectrum

the flame a vivid green 1 The green flame of botic acid, an alkyl borate, or boron fluoride gives a beautiful band spectrum in which three prominent

bands are easily recognised, the maxima being situated at  $\lambda$  5480,  $\lambda$  5190, and  $\lambda$  4930 (fig. 3).<sup>2</sup>

Silver intrate gives a white precipitate of silver metaborate with a concentrated solution of boras, and a brown precipitate of silver oxide with a dilute solution

The colour of turmeric paper, when dipped into a solution of borne acid (or of a borate acidified with hydrochloric acid) and then dired at 100°, is changed to a characteristic reddish brown (II. Rose), or, with mere traces of bonic acid, pink. In using this delicate test for bonc and it must be remembered that zucome, tantalic, columbic, and molybdic acids produce a similar coloration test is rendered more delicate by the presence of ovalic acid, the roddish-brown colour changes to a greenish- or bluish-black when sodium hydroxide is added. Borie acid turns tineture of mimosa flowers yellow, and this is in turn changed to a brick-red colour when sodium carbonate is added 5

Boric acid is neutral to litmus and methyl orange, and hence the metal present in an alkali borate may be titrated with acid as if it were present as alkalı hydroxide 6

Boric acid may be easily and accurately titrated as a monobasic acid, thus

$$H_3BO_3 + NaOH = NaBO_4 + 2H_2O_5$$

m the presence of excess of glycerol (30 per cent. by volume) or manifol (0.1 gram per cubic centi-metro of solution). The latter is the more convement to use, owing to the slight acidity of commercial glycerol, it is necessary to carry out a blank experiment when that substance is employed.

<sup>1</sup> Arfvedson's test. See Ross, Chem News, 1883, \* Alivedon's test. See Ross, Them News, 1883, 47, 186; Kunmenc, Zulsch. anal. Chem., 1873, 12, 375; Chapman, Chem. News, 1877, 35, 36; C lo Nove Foster, ibid., 1877, 35, 127, Spundler, Chem. Zeit, 1905, 29, 566, 582; Castelliana, Mti. R. Accad. Linces, 1905, [v.], 14, 1, 465, Chazzella, 1906, 36, 1, 232, Lenher and Wells, J. Amer. Chem. Suc., 1899, 21, 417.

2 See p. 12 for references.

3 Cassal and Gerrans. Chem. News, 1908, 37, 27

See p. 12 for references.
 Cassal and Gerraus, Chem News, 1903, 87, 27.
 For the turmere paper test, see Kulisch, Zeitsch angew.
 Chem., 1894, 7, 147; Lenher and Wells, loc. cit., W. H. Low, vide infra; Bertrand and Agulhon, vide infra; Jay, vide infra; Price and Ingersoll, Bull. U.S. Dept Agric. (Chem.), 1912, 137, 115; Halphen, Ann. Falsif., 1915, 8, 1.
 Robin, Compt. rend., 1904, 138, 1046; Bull. Soc. chim., 1013 [v1 v2 60?

<sup>1913, [</sup>v], 13, 602 Ann. Chim. Phys., 1830, 40, 398; Joly, Compt. rend., 1885, 100, 108; 6 Gay Lussac, Rimbsch, Ber., 1893, 26, 164.

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Alkali hydroxide free from carbonate should be used for the titration, and phenolphthalem must be used as indicator. Prior to the addition of the glycerol or mannitol, the solution, which should be slightly acid, is made accurately neutral to methyl orange or paramtrophenol and freed from carbon dioxide by boiling for a few minutes, a reflux condenser being employed to prevent loss of boric acid. (or better, the solution being boiled under diminished Mannitol or glycerol is then added and the titration of the borio acid carried out. Mineral borates soluble in hydrochlonic acid may be readily analysed by the method here outlined.2 Bone and may be titrated with alkalı hydroxide without adding either glycerol or mannitol, if tropæolin.O (i.e. sodium para-benzene-azo-resoremol-sulphonate) is used as indicator.3

Boron is determined in minerals (e.g. tournaline) by fusing with an excess of fusion mixture, acidifying with hydrochloric acid, and titrating the boric acid produced by the procedure already outlined. It is, however, first necessary to separate the boric and from the other substances piesent. This is effected most conveniently by adding a large excess of anhydrous culcium chloride, a little pure methyl alcohol, and heating to boiling while a stream of methyl alcohol vapour is blown through the liquid. The boron is volatilised as trimethyl bolate, and, together with an excess of alcohol, is condensed in a suitable receiver. A decided excess of sodium hydroxido is then added to the distillate and the alcohol distilled from the liquid, when an aqueous solution of sodium borate remains 1

<sup>1</sup> An alternative procedure consists in channating carbon dioxide from the slightly acid

<sup>1</sup> An alternative procedure consists in chammating carbon dioxide from the slightly acid (hydrochloric) solution, adding a slight excess of potassium reduce-redate mixture, and bleaching the liberated redine with sodium throughpate. In this case metals which are precipitated as hydroxides by the rodde codate mixture must be absent.

2 R. T. Thomson, J. Soc. Chem. Ind., 1893, 12, 432, Schark, ibid., 1904, 23, 699; Hoing and Spitz, Zatisch. angew. Chem., 1896, 9, 549. J. C. Jones, Imer. J. Sci., 1899, Iv. J., 7, 117; 8, 127, Copaux, Compt. end., 1898, 127, 756, Stock, ibid., 1900, 130, 516; Copaux and Boiteau, Bull. Soc. chim., 1909, [vr.], 5, 217; Saigent, J. Amer. Chem. Soc., 1899, 21, 858, W. H. Low, ibid., 1906, 28, 807, Wherry and Chapin, ibid., 1908, 30, 1687; Lindgren, ibid., 1915, 37, 1137, Mandelbaum, Zeitsch. anory. Chem., 1909, 62, 361. H. Biltz and Marcus, ibid., 1915, 77, 131, Binet du. Jassonnerv, Inn. Chem. Phys., 1909, [vin.], 17, 145. There is an extensive literature dealing with the estimation of boric acid in foods: sec, e.g., R. T. Thomson, Another, 1896, 21, 64. Cribb and Atnaud, ibid., 1906, 31, 147; Shrewshim, ibid., 1907, 32, 5. Richardson and Wallon, ibid., 1913, 38, 140, Manning and Lang, J. Soc. Chem. Ind., 1907, 26, 803, Albert and Tankaid, Pharm. J., 1904, [iv.], 19, 212, W. H. Low, loc. cit. For the detection of traces of bonax in organic matter, see Bertrand and Agulhon, Bull. Soc. chem., 1910, [iv.], 7, 90, 125, Compt. rend., 1913, 157, 1433; Jay and Dupasquer, ibid., 1895, 121, 200, Jay, ibid., 1914, 158, 357; Filippi, Ann. chim. applicating 1914, 1, 564

3 Prideaux, Zeitsch. anorg. Chem., 1913, 83, 362.

4 Whenty and Chapin, loc. ct., Stock, loc. ct., Copaux and Boiteau, loc. ct.; Sargent, loc. cit.; W. H. Low, loc. ort., 1817, 1918, 26, 21). The gravinetire estimation of boion is difficult and tedous, and olders no advantages over the volumetic estimation. The only reliable method is to hydiolyse the methyl borate, separated by the distillation process, with a known excess of pur

the volumetric estimation. The only ichable method is to hydrolyse the methyl borate, separated by the distillation process, with a known excess of pure lime, carefully evaporate, ignite, and determine the increase in weight of the lime, which is due to boron sesqui-oxide. This is Gooch's method, the lime may be replaced by sodium tangstate according to Gooch and L. C. Jones (Amer. J. Sci., 1899, [iv.], 7, 34). The distillation of the methyl borate must be so effected as to prevent the formation of dimethyl sulphate or methyl chloride. For details, see Googh, loc. cit., Gooch and L. C. Jones, loc. cit.; Moissan, Compt. rend., 1893, 116, 1087, Penfield, Amer. J. Sci., 1887, 34, 222; Arndt, Chem. Zeit., 1909, 33, 725. For the estimation of boron as boron phosphate, see Mylius and Meusser, Ber., 1904, 37, 397; as potassium borofluoride (the old method of estimation), see Stromeyer, Annalen, 1856, 100, 82°, Thaddeelf, Zeitsch. anal. Chem., 1897, 36, 568;

Perborates, when shaken with potassium dichromate solution and ether, give rise to the blue colour of perchronic acid. They may be analysed for available oxygen by titration with potassium permanganate in acid solution and by other methods.<sup>2</sup>

#### ADDENDUM

Boroethane,  $B_2H_n$ —The action of chlorine and bromine on this hydride of boron has been studied by Stock, Kuss, and Priess (Ber., 1914, 47, 3115). Chlorine causes explosions at ordinary temperatures, but bromine reacts slowly, even in the light, at 100°, however, the change is complete in a few hours. One half of the halogen used up is recovered as halogen hydride, i.e. the reaction is one of substitution, and boroethane behaves as a saturated compound. The maximum valency of boron towards hydrogen is accordingly four.

Excess of halogen leads to the production of the unmolecular haldes,  $BX_3$ , and not to  $B_2X_6$ . In order to trace the mechanism of these changes, the action of the halogens on excess of the boron hydride was studied and the following conclusions reached — The initial products are such products as  $B_2H_4X_2$  and  $B_2H_3X_3$ , which speedily decompose, without the elimination of halogen hydride, yielding  $B_2H_5X$  and  $B_2H_5$  on the one hand, and  $BX_5$  on the other. Thus, of the various halogenated derivatives theoretically possible, a mixture of the extremes is produced. No evidence of the existence of

mixture of the extremes is produced. No evidence of the existence of BHX<sub>2</sub>, B<sub>2</sub>HX<sub>5</sub>, and B<sub>2</sub>H<sub>2</sub>X<sub>4</sub> could be obtained. The monochlorade, B<sub>2</sub>H<sub>2</sub>Cl, is a spontaneously inflammable gas. The monobromide, B<sub>2</sub>H<sub>2</sub>Br, is a colourless gas with an initiating odour. It melts

at 104° and boils at c. 10 C The vapour pressure is as follows :-

Temp. °C 
$$-80^{\circ}-70^{\circ}-60^{\circ}-50^{\circ}-40^{\circ}-30^{\circ}-20^{\circ}-10^{\circ}-5^{\circ}$$
 Vap. press. in nims.  $3$  9 16 30 53 95 162 255 335

It burns with a pale green flame and fumes in the air owing to its reaction with water:-

$$B_2H_5Br + 3H_9O = B_3O_9 + HBr + 5H_9$$

It immediately reacts with potassium hydroxide to form the hypoborate KOBH<sub>3</sub>. It does not react with sodium, however, to any extent, so that the hydride B<sub>4</sub>H<sub>10</sub> cannot be thus produced.

as borne acid, by extraction with other, see Parthell and Rose, Arch Pharm, 1904, 242, 477. For a bibliography and a critical examination of various processes that have been recommended from time to time, see Sargent, low. cit., of Reischle, Zeitsch. anorg. Chem., 1898. 4, 111

<sup>1898, 4, 111</sup>Lenz and Richter, Zeitsch. anal Chem., 1911, 50, 537; Riesenfeld and Mau, Ber., 1911, 44, 3589; Bosshard and Zwicky, Zeitsch. angew Chem., 1912, 25, 938, 993

For which, see Fairar, J. Sur Dyers, 1910, 26, 81, Rupp and Mielck, Arch. Phurm., 1907, 245, 6; Lenz and Richter, loc. cit.; Latterscheid and Guggian, Chem. Zeit., 1913, 37, 200



#### CHAPTER III.

#### ALUMINIUM.

Symbol, Al Atomic weight, 27 1 (0 == 16)

Occurrence .- Aluminum is the most abundant and the most widely distributed of all the metals (Vol. 1 Chap 1), but it is nover found in the free state Among the more simple minerals in which aluminium occurs are the

Corundum . Diasporo .	$\frac{\text{Al}_2\text{O}_3}{\text{Al}_3\text{O}_4}$ $\frac{\text{H}_2\text{O}}{\text{O}_4}$	Gahnite . Cryolite .	$Z$ n Al $_2$ O $_1$ Na $_3$ Al $F_6$	•
Hydrargillite	$\Lambda I_{\bullet}O$ , $311$ , $O$	Websterite	Al,O,.SO,,911,0	
Chrysoberyl	BeALO <sub>1</sub>	Alunogen	Al <sub>2</sub> (SO <sub>4</sub> ), 1811,0	
Spinel	MgAl <sub>s</sub> O,	Alumite	Al.(SO.), 4Al(OH)K.S	0.

Bauxite is a hydrated oxide of aluminium containing more or less ferric oxide, turquoise, a hydrated aluminium phosphate

Aluminium is an essential constituent of innumerable silicates, which form the basis of most rocks and clays—Of the silicates, orthoclase, KAlSi<sub>8</sub>O<sub>8</sub>, is the most important, as it forms the chief constituent of granite, syenite, gnoiss, etc. By the weathering of orthoclase, the mineral kaolin, Al<sub>2</sub>Si<sub>2</sub>O<sub>7</sub> 2H<sub>2</sub>O, is produced. Other important silicates, such as the garnets, micas, topa:, tour maline, etc., are mentioned later (p. 92)

Aluminium is only found in small quantity in most plants, but the ashes of certain cryptogams, e.g. lycopodeum, contain large quantities of aluminium.

Aluminium is contained in the atmosphere of the sun
History.—The term "alumen" was applied by the Romans to all bodies
of an astringent taste, and among them alum was included. Alum was well known to Geber and the later alchemists, who erroneously classed it with the vitriols. This error was concerted by Paracelsus. The earth present in . alum was for a long time supposed to be calcareous In 1746, Pott stated that the basis of alum is an aigillaccons earth, and in 1754, Marggraf showed that alumina and lime are two quite distinct earths, and that alumina is present in clay, combined with silica

By the early years of the nineteenth century alumina was regarded as the

¹ Yoshida, Trans Chem. Noc., 1887, 51, 748. L'Hote, Compt. rend., 1887, 104, 863; ¹ Demarçay, ibid, 1900, 130, 91, Berthelot and André, ibid., 1895, 120, 288; Kratzmann, Pharm. Post, 1914, 47, 101, 105, Langworthy and Poter, The Occurrence of Aluminium in Vegetable, Animal Products, and Natural Waters (Wiley & Sons, 1901, Czapek, \*\*) Biochemie der Pflanien (Jena, 1905), vol. 11. p. 855; H. G. Smith, Chem. News, 1903, \*\*

oxide of an unknown metal, which H. Davy unsuccessfully endeavoured to isolate. Alummum was isolated by Wohler in 1827.

Preparation.—The oxide, sulphide, and halides of aluminium are very stable compounds, and the preparation of the metal by chemical processes is somewhat difficult From thermochemical considerations, it may be deduced that the most successful methods would be those in which an aluminium halide is reduced by fusion with an alkali metal.

It is possible that aluminium was obtained by Oersted in 1842,1 but it is generally acknowledged that the ment of having first prepared aluminium and studied its properties belong to Wohler, who, by heating anhydrous aluminium chloride with potassium, obtained the metal first in the form of a grey powder which became brilliant when burnished, and subsequently as fused metallic globules -

In 1854 both Bunsen and Deville succeeded in preparing aluminium by electrolysing the fused anhydron chloride, and in the same year Deville commenced his classic work on the manufacture of aluminium by reducing aluminium sodium chloride with metallic sodium. The following year saw the introduction of another process, namely, the reduction of cryolite by fusion with sodium
by Rose in Germany
Later, Grabeau reduced anhydrous alummum fluoride by means of sodium, and obtained aluminium of remarkable purity 5

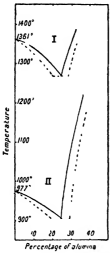
At the present time the only method by which aluminium is produced on a commercial scale is an electrolytic method, developed independently and almost simultaneously in 1886 by C M Hall in America and P V. Héroult in France. Alumna is dissolved in molten cryolite and electrolysed, when aluminium separates out at the cathode. Attempts have been made to electrolyse aluminium sulphide dissolved in fused sodium sulphide, but they have not yet been commercially successful 7

The necessity for finding a solvent for the alumina arises from the fact that its melting-point is 2010° to 2050'. The inclining-point of cryolite is a little below 1000°. In order to produce a more fluid bath, other fluorides are added In order to produce a more fluid bath, other fluorides are added to the cryolite, e.g aluminium fluoride and calcium floride

<sup>1</sup> See Berzelius, Jahresber., 1827, 6, 118; Octoted, Overs K. Danske Vulensk. Selbsk.

See Berzelius, Jahresber., 1827, 6, 118; Oorsted, Overs K. Dunske Vidensk. Selbsk. Forhandl, 1824-1825
 Wohler, Pogy Annalen, 1827, 11, 146, Ann Chim Phys., 1828, 37, 66; Annalen, 1836, 17, 43; 1845, 53, 422
 Bunsen, Pogy. Annalen, 1851, 92, 648, Doville, Ann. Chim. Phys., 1855, [in.], 43, 5; 1856, [in.], 46, 416
 Rose, Pogy. Annalen, 1855, 96, 152, Ann. Chim. Phys., 1855, [in.], 45, 369, and see also Phil. May, 1855, [iv.], 10, 364.
 Graboau, Zeitsch. angew. Chem., 1888, 1, 708, 1889, 2, 149
 Hall, US Pat., 400,664, 400,665, 400,666, 400,667; 400,766, Hénoult, Fr. Pat., 175,711 (1886), see also J. Ind. Eng. Chem., 1911, 3, 143. The following papers may also be referred to J. W. Richards, Electrochem. Ind., 1903, 1, 158, Zeitsch. Elektrochem., 1985, 1, 367; Haber, bid., 1902, 8, 607, 1903, 9, 360, Haber and Gepett, bid., 1902, 8, 126; Neumann and Olsen, bid., 1910, 16, 230, Thompson, Electrochem. Ind., 1909, 7, 19, Richardson, Trans. Amer. Electrochem. Soc., 1911, 19, 169; Fedotéev and Hjinsky, Zeitsch. angrag. Chem., 1913, 80, 113, Bock, Zeitsch. angraw. Chem., 1909, 22, 1309; Puschin, Dischler, and Maksimenko, J. Russ. Phys. Chem. Soc., 1911, 19, 169; Fedotéev and Hjinsky, Zeitsch. angrag. Chem., 1913, 80, 113, Bock, Zeitsch. angraw. Chem., 1909, 22, 1309; Puschin, Dischler, and Maksimenko, J. Russ. Phys. Chem. Soc., 1914, 46, 1347, also such text-books as Allmand, Applied Electrochemistry (Arnold, 1912); Stansheld, The Electric Furnace (M'Graw-Hill Book Co., 1914); Dony Hénault, Gall, and G'uye, Principes et applications de Pelectrochemistry (Longmans & Co., 1911), J. W. Richards, Aluminium (Sampson Low, Marston & Co., 3rd ed., 1896); Minet, L'aluminium (Pans); Thorpe, Dictionary of Applied Chemistry (Longmans & Co., 1911), vol. 1.

The equilibrium diagrams for the systems (i.) alumina—cryolite, (ii.) alumina—fluorspar, (iu.) cryolite—fluorspar, (iv.) cryolite—alumina—fluorspar, and (v.) aluminium fluorule—sodium fluorule have been partly or wholly determined. The first three systems have been studied by Pascal and Jouniaux, whose results are shown graphically in figs. 4 and 5. In each system, two series of mixed crystals are formed, and the "liquidus" curve consists of two branches meeting at a entertie point. The ternary system as worked out by the same experimenters is shown graphically in figs. 6 and 7, which represent the "liquidus" and "solidus" respectively. The



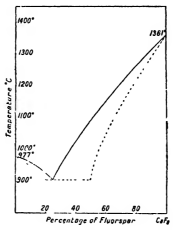


Fig. 4.—Equilibrium diagrams for the systems (I.) fluorspar - alumina, (II.) cryolite—alumina

Fig. 5.—Equilibrium diagram for the system cryolite—fluorspar.

isothermals are given for a series of temperatures separated by intervals of 50°. The ternary entectic point is 868° and corresponds to a mixture of the following composition eryolite, 59 3 per cent; fluorispir, 23 0 per cent; alumina, 17·7 per cent. In actual practice, according to Pascal and Jounnaux, alumina is added so as to constitute 10 to 25 per cent, of the alumina-cryolite mixture, and fluorispar is added up to 36 per cent, of the weight of the cryolite present. The working compositions are therefore included in the trapezium bounded by the lines Fb, Fc, bc, and  $\Delta a$ . This trapezium includes the entectic mixture, and no mixture in it has a melting-point higher than 980°. The temperature limits adopted in practice are given by Pascal and Jounnaux as 875° to 950°.

The system aluminium fluoride --sodium fluoride has been studied by

Pascal and Jouniaux, Bull. Noc. chim., 1913, [iv], 13, 439, Rev. Mctallurgie, 1914, 11, 1069; cf. Pyno, Trans. Amer. Electrochem. Soc., 1906, 10, 63, Moldenhauer, Metallurgie, 1909, 6, 14; Lorenz, Jabs, and Eitel, Zeutsch. anorg. Chem., 1913, 83, 39.

1.00

Fedotéev and Hjinsky, whose results are shown graphically in fig. 8. The branches AB, BCD, and DE correspond to the solid phases NaF, 3NaF.AlF<sub>3</sub>, and 5NaF.3AlF<sub>3</sub>, respectively, and the latter compound breaks up at 725° into 3NaF.AlF<sub>3</sub>, and AlF<sub>3</sub>. It will be seen that the addition of aluminium fluoride to cryolite produces a readily fusible mixture. Fedoteev and Hjinsky recommend that the solvent for alumina should be prepared by adding sufficient aluminium fluoride to cryolite to give the mixture the composition G (fig. 8); that the temperature should be maintained at 900°, and that the quantity of alumina added should not exceed 7.5 per cent of the weight of the solvent. If pure cryolite be used as solvent, Fedotéev and Hjinsky recommend that not more than 10 per cent of alumina be added

It is essential that the molten metal shall be liberated at the cathode in

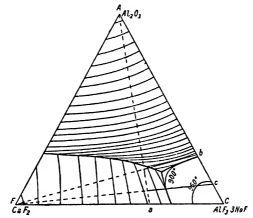


Fig. 6.—The system ciyolite—alumina—fluoispar Projection of "liquidus." Contour lines for each 50° difference in temperature.

a medium of lower specific gravity than the metal itself. At the ordinary temperature, aluminium is less dense than cryolite, fortunately, however, the reverse is true at high temperatures, as will be seen from the results tabulated below and represented diagrammatically in fig.  $9...^2$ 

```
658°
                                           6829
                                                   740
                                                                           925°
                                                                                   10009
Temperature
                                                  2 43
                                                                           2.37
                                                                                   2.35
                                          2 45
                                                                   2:39
                                 . 246
Density of Al (liquid)
                                 . 972°
                                          975
                                                                   995°
                                                                           1003
                                                                                   1018
                                                                                          1032°
Temperature
                                                                  2·220
1075°
                                          2·197
1048°
                                                  2 203
1061°
                                                          2·197
1069°
                                                                                  2·196
1083°
Density of cryolite (liquid)
                                . 2·185
                                                                           2 209
                                                                                           2.189
                                                                           1070
Density of cryolite (liquid)
                                  2.177
                                          2.167
                                                  2 154
                                                          2 139
                                                                  2.129
```

It will be seen that molton cryolite, like water, has a point of maximum density, the density then being 2.22 and the temperature 995. The density of cryolite is diminished by the addition of silica and increased by the addition of calcium fluoride (see fig. 9); the influence of alumina on the

<sup>&</sup>lt;sup>1</sup> Fedotéev and Iljinsky, loc cit; see also Lorenz, Jabs, and Eitel, Zeitsch. anorg Chem., 1913, 83, 39, 328.

5 Pascal and Jounnaux, Bull. Soc. chim., 1914, [iv.], 15, 312.

density is rather complex, but unless a large quantity (over 20 per cent.) is added, the density is diminished (see fig. 9). The densities of the commercially important ternary mixtures of cryolite, alumina, and fluorspar at 950° are shown graphically in fig. 10, it will be seen that they are less than 2.40, and in actual practice the presence of a little silica in the cryolite makes them all a little lower than the values given in the diagram.

The Hall and Héroult processes only differ slightly in the nature of the plant employed, and perhaps to some extent in the composition of the electrolyte adopted. Each Hall cell consists of a rectangular cast-non box,  $6' \times 3' - 3'$ , thickly lined with carbon. This lining forms the cathode. The actual depth of electrolyte in the bath is 6 inches. As anode, a large number of carbon rods are used, which dip into the electrolyte and end about 1

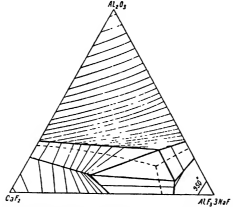


Fig. 7 —The system cryolite—alumina—fluoispar Projection of "solidus." Contour lines for each 50° difference of temperature.

mch above the level of the aluminium at the bottom. It is therefore clear why the density of the bath must be such as to allow the aluminium, as it is produced, to sink rapidly to the bottom, otherwise, besides oxidation of the metal taking place, short circuiting will occur. About 15 to 20 parts of alumina are added for each 100 parts of solvent. The electrolyte is roughly covered with a layer of carbon, on the top of which alumina is placed. As electrolysis proceeds, this alumina is stirred into the bath and at intervals the separated aluminium is tapped off from the bottom.

The current consumed by each cell is 10,000 ampères at 5 5 volts, the current density being 100 amp/dem.² at the cathode, and about 500 amp./dem.² at the anode. The current efficiency does not exceed 70 per cent, largely owing to the formation of a "metal mist" of aluminium particles which diffuse to the surface and become oxidised. The least potential difference capable of producing continuous electrolysis is 2·1 to 2·2 volts, which value does not differ greatly from the approximate figure calculated from the

Prepared as described later (p 79).
Clacher, Met. Chem. Eng., 1911, 9, 137.

heat of formation of alumina. The potential differences required for the electrolysis of aluminium chloride, aluminium fluoride, sodium fluoride, and calcium fluoride are approximately 23, 40, 47, and 47 volts respectively.

The cost of the carbon anodes, which are eaten away by the liberated oxygen with the formation mainly of carbon monoxide, is a very serious item. The anode rods must be evenly hard, very slightly porous, and leave very little ash when burnt They are usually made from petroleum coke 1 Waterpower is always employed in the production of the necessary electric current. The world's supply of aluminium is produced mainly by the Aluminium Company of America, at Niagara, Massena, and Shawingian Falls, the Société Electrométallurgique Française, at Froges, La Plaz, and St Michael, and the British Aluminium Company, at Kimochleven in Scotland The world's output of aluminium was about 8000 tons per annum from 1900-1905; since

then it has steadily increased, and was about 30,000 tons in 1909.2

.. . . .

Aluminium produced by electrolysis contains 99 per cent. or more of alumnium, the chief impurities being non and silicon A little carbon is also present, and m a sample exammed spectroscopically by Hartley and Ramage,3 traces of sodium, potassium, calcium, copper, silver, manganese, lead, gallium, and indium were detected. It is extremely difficult to purify the aluminium after it has once been produced, and hence it is necessary to employ pure materials in its preparation. It is for this reason that the carbon anodes must be practically free from ash. The alumina is usually prepared from bauxite by a method described later (p 79). The cryolite is not specially

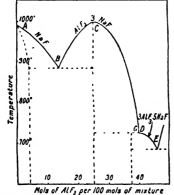


Fig 8 .- Equilibrium diagram for the system sodium fluoride-alum mum fluoride.

purified, but any foreign metals present are removed after it has been submitted to the action of the current for a short time.

Pure aluminium is best prepared by reducing pure, redistilled aluminium · tribromide with a slight deficiency of sodium. The tribromide is mixed with sodium and potassium chlorides and heated with metallic sodium in a crucible lined with a mixture of alumina and sodium aluminate.1

Properties.—Aluminium is a tin-white metal which may be highly The commercial metal is extremely sonorous,5 mallcable, and ductile. It is best worked at 100° to 150°, at 600° it is easily broken, and at a slightly ligher temperature it may be powdered in a mortar.

<sup>1</sup> Clachet, Met Chem. Eng., 1911, 9, 137.

1 Thorpe, A Inctionary of Applied Chemistry (Longmans & Co., 2nd ed., 1911-1913), vol. 1. p. 106.

2 Hartley and Ramage, Trans. Chem. Soc., 1897, 71, 547.

4 Mallet, Phil. Trans., 1880, 171, 1022; Chem. News, 1882, 46, 178.

5 The pure metal is not sonorous (Mallet, cited by Baskerville, J. Ind. Eng. Chem., 1814, 6, 183).

<sup>1914, 6, 183).</sup>Granger, Bull. Soc. chim., 1902, [mi.], 27, 789.

Aluminium has a crystalline structure when slowly cooled. Apparently the crystals are regular octahedra.¹ After being worked, the metal is devoid of

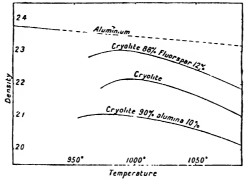


Fig. 9.—Densities of aluminium, etc., at high temperatures

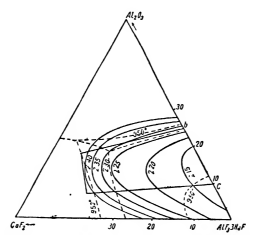


Fig. 10 -Densities of mixtures of civolite, fluorspar, and alumina at 950°.

structure. Cast alummium is about as hard as silver, and becomes harder when hammered.2

The density of aluminum varies with the treatment to which it has been subjected. The density of the cast metal (containing 0.36 per cent,

Deville, Ann Chim. Phys., 1855, [in ]. 43, 11
 For the manufacture of aluminium foil and powder, see Guillet, Rev. Mitallurgie, 1912, 9, 147.

impurities) is 2.703 at 18° C.; that of the worked metal may be less than 2.703, but it increases on annealing and may reach the value 2.7085 at 18°.1 Mallet 2 gives the density of pure aluminium as 2.583 at 4°C. The mean coefficient of expansion between 0° and 100° is  $2.432 \times 10^{-5}$  for hard-drawn metal and 2.454 10-5 for the annealed metal 3

The value of Young's modulus for aluminium is about 6 91 103 (in kilos. per sq. mm) at 20°, the rigidity being about 2.7 103 (in kilos. per sq. mm), both these values diminish considerably with rise of temperature. The compressibility is  $1.49 \times 10^{-6}$  per atmosphere. The tensile strength is affected by the form, method of casting, and subsequent treatment, the ultimate strength being (in tons per sq inch) 7 in castings, 11 in sheet aluminium, and from 13 to 29 in aluminium

The thermal conductivity of aluminium is 0.3435 at 0° and 0.3619 at 100° (absolute units), according to the measurements of Lorenz, while Jaeger and Disselhoist give 0 1801 and 0:4923 as the corresponding figures for aluminium containing 0.4 per cent of copper and 0.5 per cent of non. Aluminium is therefore a slightly better heat conductor than zinc. The electrical conductivity is 35.6 104 reciprocal ohms per cm cube at 0°, according to Sturm, while the conductivity at 0° and temperature-coefficient of the resistance (between 0° and 100°) are given by Broniewski as 40 1 104 and  $4.25 \times 10^{-3}$  respectively for annealed aluminium, and 38.5 10<sup>4</sup> and 4.10 × 10-3 for the chilled metal 8

Aluminium melts at 658 ± 1°,9 boils at 1800°,10 and expands at 4.8 per cent. on fusion. 11 Owing to the formation of a protective film of oxide, it is possible to heat a piece of aluminium wire above the melting-point without destroying its shape 12 Being a metal of low atomic weight, aluminium has a high specific heat, which, moreover, has a large temperature-coefficient. According to E. H Gushiths and E Gushiths, the specific heat is 0 2096 at 0° and 0 2252 at 100° 13 Intermediate values are given by the expression  $s = 0.20957(1+9\ 161\ 10^{-1}t-1\ 7\ 10^{-6}t^2)$ , and hence the mean specific heat is 0.2180 between 0° and 100°, and 0.2196 between 20° and 100°. For the latter value, Schmitz found 0 2191 14 Bontscheff has experimented over a 

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<sup>1</sup> Brislee, Trans. Fanaday Soc , 1912, 7, 221; 1913, 9, 162, Lowiy and Paiker, Trans.
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<sup>Brislee, Trans. Kanaday Soc., 1912, 7, 221; 1913, 9, 102, Lowiy and Paiker, Trans. Chem. Soc., 1915, 107, 1005
Maillet, them New, 1882, 46, 178
Brislee, tw. cit., and them New, 1912, 105, 3, cf. Fizeau, Compt. rend., 1869, 68, 1125; Le Chatcher, thid., 1889, 108, 1096.
Brislee, Trans. Faraday Soc., 1913, 9, 155; Voigt, Wied Annalen, 1893, 48, 674, G. S. Moyer, thid., 1896, 59, 668, Schaefer, Ann. Physik, 1901, 5, 220, 1902, 9, 665, 1124; Pionchou, Compt. rend., 1892, 115, 162., Slotte, Acta Soc. Sci. Fennice, Helsingfors, 1800, 56, 1910, 20.</sup> 1124; Pionehon, Compt. v. nd., 1892, 115, 162. Slotte, Acta Soc. Sci. Fernice, Helsingfors, 1899, 26; 1900, 29

8 Richards (and others), J. Amer. Chem. Soc., 1909, 31, 154; 1915, 37, 164.

6 L. Lorenz, Wied. Annalen, 1881, 13, 422, 582

7 Jaeger and Disselhoust, Wies. Abh. phys. tech. Reichsunstall, 1900, 3, 269

8 Sturm, Innaquard Dissertation (Rostock, 1904); Bioniewski, Ann. Chim. Phys., 1912, [viii.], 25, 5; cf. J. W. Richards, Chem. News, 1897, 75, 217.

10 Day, Sosman, and Allen, Amer. J. Sci., 1910, [iv.], 29, 93; Burgess, J. Washington. Acad. Sci., 1911, 1, 16; Chem. News, 1911, 104, 165.

10 Greenwood, Proc. Roy. Soc., 1909, A, 82, 396, 1910, A, 83, 483.

11 Toepler, Wied. Annalen, 1894, 53, 343.

12 Matignon, Bull. Soc. chim., 1909, [iv.], 5, 91; Moniteur Sci., 1900, [iv.], 14, 357; Von Bolton, Zeitsch. Elekthochem., 1908, 14, 766.

10 E. H. Griffiths and E. Griffiths, Phil. Trans., 1913, A, 213, 119.

ALUMINIUM. 55.

wide temperature interval and gives the following values for the specific heat :--

Temp. °C. . 650° - 100° O' 100° 300 500° 0.2739Specific heat 0 1893 0 2089 0 2226 0 2434 0.3200

The results may be expressed by the formula -

$$s = 0.20890 + 1.6187t \times 10^{-4} - 2.9125t^2 \cdot 10^{-7} + 4.6183t^3 - 10^{-10}$$

At -193.9° the specific heat of aluminium is only 0.086, and at -253.9it has fallen to 0 0024, at these extremely low temperatures the specific heat is proportional to the cube of the absolute temperature 2 The atomic heat of aluminium at ordinary temperatures is rather lower than might be anticipated, since, adopting the mean specific heat between 20° and 100°, the atomic heat is only 5.95°. Girlliths and E. Girlliths give the following as the most probable values of the atomic heat of aluminium at various temperatures -- 4

Temp (abs.) . 32 4° 80° 120° 200′ 250 300' 310" 380°  $0.25 \quad 2.27 \quad 3.74 \quad 5.14 \quad 5.54$ 581 598 613 Atomic heat

The latent heat of fusion of aluminium is 70 to 80 cals, per gram 5

Aluminium is paramagnetic, the magnetic susceptibility at ordinary temperature being +18×10-6 cgs. electromagnetic units per unit volume. The atomic refraction of aluminium in its compounds is 9 5 (for the Ha line; Gladstone and Dale's formula) 7

The electrode potential of aluminium is not known with any accuracy, the value  $\epsilon_n=-103$  to 1.28 volts being uncertain and probably numerically too great—the potential series aluminum probably occupies the following position -

Alkah and alk, earth metals, Mg, Mn, Al, Zn, Cr, Cd, Fe, etc 8

When aluminium is used as anode in passing a current through an aqueous solution, and the voltage does not exceed some 25 volts, the current that passes quickly falls to almost zero. This seems to be due to the formation of a layer of aluminium hydroxide on the anode. The critical voltage, above which an appreciable current can be made to flow, varies with the

<sup>&</sup>lt;sup>1</sup> Bontscheff, Inauquial Dissertation (Zurich, 1899), also in Laemmel, Ann. Physik, 1905, [1v], 16, 551

Neinst and Schwers, Nizanayber K. Akad. Wess Berlin, 1914, p. 355
 For other work on the specific heat of aluminum, see Pronchon, Compt. rend., 1892, 175, 162; Tilden, Phil. Trans, 1900, A. 194, 233, 1903, A. 201, 37, Trans Chem. Soc., 1905, 18, 551; Trowbridge, Searne, 1808, 8, 6; Behn, Wad. Annalen, 1808, 66, 237; Schubel, Zeitsch, amorg Chem., 1914, 87, 81, and Vol. I., Chap. II
 E. H. Griffiths and E. Griffiths, Phil. Trans, 1914, A. 214, 319.
 Pronchon, lowed; Laschtschenko, J. Russ. Phys. Chem. Soc., 1914, 46, 311; cf.
 J. W. Richards, Aluminium (Band. & Co., 3rd. ed., 1896), p. 80
 Wills, Phil. Mag., 1898, [v.], 45, 432; Koenigsberger, Wed. Annalen, 1898, 66, 688; Honda, Ann. Physik, 1910; [v.], 32, 1027, Owen, and., 1912, 37, 657, Proc. K. Akad. Wetensch. Amsterdam, 1911, 14, 627
 Gladstone, Proc. Roy. Soc., 1807, 60, 140
 Neumann, Zeitsch. physikal. Chem., 1894, 14, 217; Wilsmoie, ibid., 1900, 35, 318; Burgess and Hambuchen, Electrochem. Ind., 1903, 7, 165; van Deventer and Lummel, Gasm. Weekblad., 1907, 4, 771; 1908, 5, 359; van Laar, ibid., 1908, 5, 124. Nernst and Schwers, Sitzungsber K. Akad. Was Berlin, 1914, p. 355

1000

temperature and the nature of the electrolyte. This property of aluminium is utilised at times for obtaining a direct from an alternating current.1

The arc and spark spectra of aluminium 2 are fairly simple. The most intense lines in these spectra, i.e. the "hauptlinien," are (Exper and Haschek) the following :-

arc: 2568 08, 2575 20, 2652 56, 2660 50, 3082 30, 3092 89, 3944 20, 3961.71 spark: 2816.41, 3082 30, 3092.89, 3944.22, 3961.74, 4529 70, 5696.71.

The most persistent lines in the spark spectrum of aluminium, ie the "ultimate" lines, which should be looked for when seeking the traces of aluminium, are (Exner and Haschek's wave-lengths) 396174,\* 394122,\* 3092.82, 3082.30, 2816.41, those asterisked being the most sensitive.

The arc spectrum of aluminium contains a number of very characteristic bands due to aluminium oxide, which disappear when the arc is surrounded by an atmosphere of hydrogen.

Aluminium always contains a little occluded gas, which may be extracted by fusing the metal in vacuo.1

Aluminium readily combines with the halogens. A compact piece of the metal is only superficially oxidised when heated in air or oxygen, but in thin foil it burns brilliantly when heated in oxygen. Aluminium powder begins to oxidise rapidly at 400°, and readily burns if strongly heated at one point, aluminium oxide and a little nitride being produced.5 Aluminium rapidly oxidises if the surface is amalgamated with mercury, an arborescent growth of alumina quickly forming all over the metal 6 At high temperatures finely divided aluminium readily unites with sulphur, selenium, phosphorus, and arsenic, it also combines with antimony, but with more difficulty.7 Aluminium combines directly with nitrogen, producing a nitride, it also combines with carbon, silicon, and boron.

Owing to the extremely large heat of formation of aluminium oxide, aluminium is able to reduce many oxides, with the evolution of much heat, Thus, aluminium powder burns readily when heated in the oxides of sulphur,

<sup>1</sup> W W. Taylor and Inghs, Phil May, 1903, [vi.], 5, 301, F. Fischer, Zeitsch. physikal. Chem., 1901, 48, 177, Zeitsch. Elektrochem, 1904, 10, 869, Zeitsch. anory. Chem., 1905, 43, 341, Charters, J. Physical Chem., 1905, 9, 110, Baristo and Meiser, Thins. Faraday Soc., 1911, 7, 1; Baristo, ibid., 1912, 8, 232; Schultze, Zeitsch. Elektrochem., 1914, 20, 307, 592.

\*\*Kayser, Handbuch der Spektroskopie (Leipzig, 1900-1912), vol. v p 94; Exner and Haschek, Die Spektren der Elemente bei normalem Druck (Leipzig and Wien, 1911); Grunter, Zeitsch. Wiss. Photochem, 1913, 13, 1 (arc and spark), Edgent, ridd., 1913, 13, 20; 1914, 14, 137 (spark, ultraviolet); Huppers, ibid., 1913, 4, 46 (arc, below 3200); de Gramont, Compt. rend., 1913, 157, 1364 (band spectrum); Stark and others, Ann Physik, 1918, [iv.], 42, 241; 45, 29 (canal ray spectrum), Pollok, Sc. Proc. Roy. Dubl. Soc., 1912, 23, 202 (vacuum tube spectrum), L. and E. Bloch, Compt. rend., 1914, 158, 1416 (extreme ultra-violet, spark).

<sup>1890.

9</sup> Jehn and Hinze, Ber., 1874, 7, 1498; Lebon, Compt. rend., 1900, 131, 706; Jourdain, 1910, 150, 391.

7 Fonzes-Diacon, 181d., 1900, 130, 1314.

nitrogen, and carbon 1 The reduction of a solid oxide is best accomplished by mixing the powdered oxide with an equivalent of granulated aluminium and starting the reaction by putting a little barrum peroxide and magnesium powder mixture on the top and lighting it with a match or a piece of burning magnesium ribbon. In this way the oxides of non, manganese, chromium, etc., can be easily reduced to metal, so much heat being generated that both the metal and the alumina produced are melted in the reaction 2. These reductions are generally known as "thermit" reactions (vide infra, p. 58). In a few cases the reduction of an oxide by aluminium is an endothermic change and can only be brought about by supplying the necessary heat, e.g. the reduction of calcium and magnesium oxides.3 Thermit reactions may be readily applied to the preparation of phosphides, arsenides, silicides, and borides by simultaneously reducing two oxides 1

At ordinary temperatures aluminium is unattacked by air-free water, but ordinary water slowly acts upon it. The rate of corrosion is greatly mercased by the presence of impurities, particularly by traces of sodium, copper, or iron, and is also augmented by rise of temperature " Aliminium powder slowly decomposes water at 100°, and the powder, when ignited in air and plunged mto steam, continues to burn, with the evolution of hydrogen 6 Alpininium amalgam decomposes water readily, aluminium hydroxide and hydrogen being produced 7. To effect this reaction it is only necessary to amalgamate superficially the surface of aluminium foil by namersing it in aqueous mercuric chlorde, and then wash the foil with cold water. The aluminium-mercury couple thus formed, owing to the ease with which it decomposes water, constitutes a valuable reducing agent \

Aluminium is rapidly corroded by dilute hydrogen peroxide, aluminium hydroxide being formed b

Aluminium dissolves readily in hydrochloric acid, dilute or concentrated, hydrogen and aluminium chloride being produced. The action of dilute sulphuric acid is very slow, the concentrated acid attacks the metal with the evolution of sulphur dioxide Phosphoric acid, dilute or concentrated, readily attacks aluminum, hydrogen being evolved. In comparison with its action on other metals, the action of intue acid on aluminum is extremely slow. With 5 to 20 per cent acid at 25' to 30' the main reaction is as follows -

# $A1 + 411 \text{NO}_3 = A1(\text{NO}_3)_3 + 211_20 + \text{NO}_4$

With a large excess of acid, introgen appears among the gaseous products. A little ammonium nitrate is produced. Nitric acid of density 1.15 dissolves aluminum faster than the 145 acid, and the rate of solution of

Mallet, Trans. Chem. See, 1876, pt. 2. 349; Franck, Bull. Soc. chim., 1894, [iii.],
 11, 439; Guntz and Masson, chil., 1897, [iii.], 17, 209. Matignon, loc. cit.
 Goldschmidt, J. Soc. Chem. Ind., 1898, 17, 543. Electrochem. Ind., 1908, 6, 360.
 Weston and Ellis, Trans. Fainday Soc., 1908, 4, 130., 1909, 4, 166. Matignon, Compt. rend., 1913, 156, 1157.
 Colani, Compt. rend., 195, 141, 33. Matignon and Trannoy, chid., 1905, 141, 190.
 Donatti, Zeitsch. ang. w. Chem., 1895, 8, 111. Bailey, J. Soc. Chem. Ind., 1913, 32, 293; Scala, Alti. R. Accod. Linear, 1913, 22, 143, 95.
 Matignon, Compt. rend., 1906, 130, 1390.
 Baille and Fery, Ann. Chim. Phys., 1889, [v.], 17, 246.
 Cohen and Ormandy, Trans. Chem. Soc., 1890, 57, 811. Wishcenus, J. prakt. Chem., 1896, [ii.], 54, 18, Lebon, Compt. end., 1900, 131, 706.
 Droste, Chem. Zest., 1913, 37, 1311.

the motal increases with the fineness of division, thick foil dissolving very slowly, but coarse turnings much more rapidly. Organic acids attack aluminium only very slowly, but if the protecting layer of hydrogen that forms on the metal is removed (e.g. by operating in a vacuum) the rate of solution is greatly accelerated 2

Alkalı hydroxides in aqueous solution rapidly dissolve aluminium, alkali aluminates and hydrogen being produced. With ammonium hydroxide, the products are aluminium hydroxide and hydrogen Aluminium also dissolves in aqueous alkalı carbonates, carbon dioxide and hydrogen being evolved. Aqueous solutions of salts, eg. sodium chloride, slowly attack aluminium, but only in the presence of oxygen; the addition of a small quantity of a weak organic acid hastens the corrosion.

Colloidal aluminium. -- When two aluminium rods are immersed in water, their ends being separated by only 0.1 mm, and a condensed spark discharge passed between them by means of a powerful induction coil, a colloidal solution of aluminium is obtained.3

Applications.—Aluminum is largely employed in the iron and steel industry, as when it is added to molten steel (a few ounces per ton) it removes the dissolved gases to a large extent, and thereby prevents the development of blowholes in castings. In America, aluminium is extensively used in the place of copper for electrical transmission. It is also employed in the production of carbon free chromium, manganese, molybdenum, ferrotitanium, ferro-vanadium, ferro-boion, manganese-copper, manganese-titanium, chromium-manganese, etc The metallic oxide or mixture of oxides is reduced by means of a slight deficit of granulated aluminium, the reactions being carried out in large magnesia-lined crucibles as described previously (p. 57). A mixture of powdered aluminium and granulated rolling-mill scale (Fe<sub>3</sub>O<sub>4</sub>), known by the registered name of thermit, is extensively used in welding operations. When fired in the usual way (p. 57) the iron oxide is rapidly reduced to metallic iron, and so much heat is developed that both the iron and alumina are produced in the liquid state. Train and railwayline sections may be joined, and broken engine-frames, driving rods, crank shafts, etc., may be repaired by igniting a charge of thermit and allowing the molten iron to flow around the place at which a join is to be effected.4

The light aluminium alloys (p. 61) containing a high percentage of duminium are largely used for constructional purposes, e., for parts of notor-cycles, motor-cars, and aeroplanes. The salts of aluminium are nonpoisonous, and hence aluminium is used to a considerable extent in the nanufacture of kitchen utensils, particularly for army use, and is coming nto use in the construction of chemical plant.<sup>5</sup> Aluminium cannot be atisfactorily soldered, despite the large number of solders that have been natented for the purpose. It must be riveted or welded.

Aluminium powder enters into the composition of certain explosives.

<sup>1</sup> Stillman, J. Amer. Chem. Soc., 1897, 19, 715 (a complete bibliography is given here); Fatson Smith, J. Soc. Chem. Ind., 1904, 23, 475; van Doventer, Chem. Weekblad., 1907, 69; Hale and Foster, J. Soc. Chem. Ind., 1915, 34, 464.

2 Intte, Compt. rend., 1898, 127, 919; 1899, 128, 195, 793.

3 Zavriev, Zeitsch. physikal. Chem., 1914, 87, 507.

4 Goldschmult, Skuhl und Eisen, 1898, 18, 408; Zeftsch. Elektrochem., 1898, 4, 494; 899, 6, 53; J. Soc. Chem. Ind., 1898, 17, 543, 584, 619; Electrochem. Ind., 1908, 6, 860; Norpe, A Dictionary of Applied Chemistry (Longmans & Co., 2nd. ed., 1912–13), vol. v.

Seligman, Science Progress, 1912, 6, 615; Mojssan, Compt. rend., 1899, 128, 895.

See Bichel, Zeitsch. angew. Chem., 1905, 18, 1889.

Thus, ammonal contains 93-97 per cent. NH<sub>1</sub>NO<sub>3</sub>, 4-6 per cent. Al, and 0-1 per cent. moisture, and is a good mining explosive and high explosive for shells <sup>1</sup> Ammonal B contains 94-96 per cent. NII<sub>1</sub>NO<sub>3</sub>, 2.5-3.5 per cent. Al, 2-3 per cent. wood charcoal, and 0-1 per cent moisture. Another aluminium explosive <sup>2</sup> contains 85.5 per cent. NII<sub>1</sub>NO<sub>4</sub>, 8 per cent. Al, 25 per cent carbon (hydrocarbon or introhydrocarbon), and 4 per cent.

K<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub>.3

Atomic and Molecular Weights .- From the analyses and molecular weights of aluminium chloride, bromide, and iodide, various double compounds that the preceding salts form with organic substances, aluminium methyl, aluminium ethyl, and aluminium acetylacetonate, it follows that the atomic weight of aluminium is approximately 27, or three times its combining weight. This conclusion is in harmony with Dulong and Petit's Law, the isomorphism of (i.) alumina, ferric and chromic oxides, (n) the aluminum, gallium, indinin, chromium, non, vanadium, cobalt, and rhodium alums, (iii) the aluminium, non, chromium, and cobalt double oxalates, etc.

The atomic weight was determined by Mallet im 1880 by four distinct methods, the results of which are appended -5

```
(\mathbf{NH_1})_2\mathbf{SO_4}.\mathbf{Al_2}(\mathbf{SO_3})_3.2441_2\mathbf{O}:\mathbf{Al_2}\mathbf{O}_4
                                                            100 11 2793 . . Al 27 152
                                   3Ag : AlBi,
3H<sub>2</sub> · 2Al
2Al · 3H<sub>2</sub>O
                                                            100 82 155 .: Al 27 109
                                                           100 , 896 33 , Al = 27 095
                                                           100 99 818
                                                                                 .: Al 27 072
```

The first ratio was determined by calcining ammonium alum and measuring the loss in weight, the second, by titrating aluminium bromide against silver according to the procedure of Stas. The third was determined by dissolving aluminium in sodium hydroxide and measuring the hydrogen evolved, and the fourth was carried out like the third, the hydrogen, however, being burnt to water. Other determinations are of less importance. The atomic weight of aluminium is at present taken as Al 271.

From the freezing point determinations made by Heycock and Noville, it appears that aluminum in dilute solution in tin has the molecular formula Al2, but it is not certain that pure tim separates on freezing. Alumnum likewise appears to be present largely as diatomic molecules in solution in mercury 8

Von Dahmen, Eng. Pat., 16,277 (1900).
 See Eng. Pat., 16,511 (1901)
 On aluminum condensers for laboratory use, see Mastbaum, Chem. Zeil., 1910, 34, 1319, for aluminum stopcocks on bydrogen sulphide apparatus, see Campbell, J. Amer. Chem. Soc., 1911, 33, 947; and on the use of aluminum dishes in quantitative analysis, see Formánek and Pet. Chem. Zeil., 1909, 33, 1282.
 Mallet, Phil. Trans., 1880, 171, 1003
 Clarke, A Recalculation of the Atomic Weights, 3rd ed. ("Smithsonian Miscellaneous Collections," vol. 54, No. 3, 1910). The anti-cedent data have been changed to the following:— O=16 000, H=1 00762, Ag-107 880, Br=79 916, S=32 065, N=14'008.
 Berzelius, Poya Annalen, 1826, 8, 177, Vather, Amer. J. Seil., 1835, 27, 241; "Tissier, Compt. rend., 1858, 46, 1105, Isnaid, ibid., 1868, 66, 503; Dumas, Ann Chim. Phys., 1858, [m], 55, 151; Terral, Bull. Sec. chim., 1879, 31, 153, Baubigny, Compt. rend., 1888, 97, 1369, Thomsen, Zeitsch. amorg. Chem., 1895, 11, 14, 1897, 15, 447; Kohn-Abrest, Bull. Soc. chim., 1905, [iii], 33, 121
 \* Heycock and Neville, Trans. Chem. Soc., 1890, 51, 376.

# <sup>₹/36</sup>6L

#### ALLOYS AND COMPOUNDS OF ALUMINIUM.

Aluminium forms one series of salts, in which it is tervalent. The salts are derived from the basic oxide  ${\rm Al_oO_3}$ . Those derived from colourless acids are themselves colourless. The soluble salts from hydrates containing considerable water of crystallisation, and the anhydrous salts, dissolve in water with the evolution of a large amount of heat. The salts are not poisonous

On the ionic hypothesis the salts of aluminium are dissociated in aqueous solution, giving rise to a colourless cation Al. :-

$$AlX_1 = Al\cdots + 3X'$$
.

The ionic mobility of the cation, \( \frac{1}{3} \) Al..., is 40 4 at 18° C.\( \frac{1}{2} \) Despite the high position occupied by aluminium in the electromotive series (p. 55), aluminium oxide is only a weak basic oxide and aluminium hydroxide but a feeble base. Accordingly, in aqueous solution, aluminium salts are appreciably hydrolysed, and those derived from moderately strong acids give a decidedly and reaction. A solution of 1 gram-molecule of aluminium chloride or brounde in 1000 litres of water is hydrolysed to the extent of 4 per cent at 25', while the percentage hydrolysis (v) of the chloride in aqueous solution at 997 varies with the dilution (" litres per gram-molecule), as follows — 2

v	32	64	128	256	512	
,	8.0	13 2	19.7	989	41.1	

Aluminium hydroxide is a stronger base than ferric hydroxide, but is weaker than beryllium hydroxide. It is possible that, owing to the transformation of aluminum hydroxide into the form of a colloidal hydrosol, the degree of hydrolysis of alummium salts gives a somewhat evaggerated idea of the weakness of the hydroxide as a base

Owing to the appreciable hydrolysis of aluminium salts, the soluble salts derived from volatile acids cannot be prepared in the anhydrous state by the evaporation of their aqueous solutions on the steam-bath

Aqueous solutions of aluminium salts dissolve an appreciable amount of aluminium hydroxide.

Thermochemistry of Aluminium Compounds.3—The heats of formation, etc, of the more important compounds of aluminium are given in the following table, in kilogram-calories -

A. Heydweller, Zeitsch physikal. Chem., 1915, 89, 281
 Ley, Zeitsch, physikal. Chem., 1901, 30, 245; Kablukoff and Sachanoff, ibid., 1909, 69,
 19; Povarnin, J. Russ. Phys. Chem. Noc., 1909, 41, 1014, 1910, 42, 207; Wood, Trans. Chem. Noc., 1908, 93, 417; van Pelt, Bull. Noc. chem. Belg., 1914. 28, 101; Kullgren, Zeitsch. physikal Chem., 1913, 85, 466.
 J. Thomsen, Thermochemische Untersuchungen (Leipzig, 4 vols., 1882-1886), Berthelot, Thermochemisc (Paris, 2 vols., 1897); Ann Chem. Phys., 1901, [vii.], 22, 479, Baud, bid., 1904, [vii.], 1, 8; T. W. Richards and Burgess, J. Amer. Chem. Soc., 1910, 32, 431;
 T. W. Richards, Rowe, and Burgess, ibid., 1910, 32, 1176, Rolla, Gazzita, 1915, 45. 1.
 192; Mixter, Amer. J. Not., 1915, [vv.], 39, 295.

#### ALUMINIUM.

Compound.	Heat of Formation.	Heat of Solution
AlF <sub>1</sub>	Cals 219 0 161 1 122 0 70 3 380 2 297 0 126 4 879 71	Cals +31 3 1 7 1 77 6 + 85 3 1 89 0  

The following data refer to heats of reaction -

```
The following data refer to heads  \begin{bmatrix} Al_2O_3 \end{bmatrix} \operatorname{cryst.} + \begin{bmatrix} Na_2O \end{bmatrix} & + & 48 & 0 & 0 \\ \begin{bmatrix} Al_2O_3 \end{bmatrix} \operatorname{amorp.} + \begin{bmatrix} Na_2O \end{bmatrix} & + & 55 & 1 \\ Al_2O_3 \end{bmatrix} & + & 55 & 1 \\ Al_2O_3 \end{bmatrix} & + & 55 & 1 \\ Al_2O_3 \end{bmatrix} & + & 127 & 0 \\ Al_2O_3 \end{bmatrix} & + & 127 & 0 \\ Al_2O_3 \end{bmatrix} & + & 127 & 0 \\ Al_2O_3 \end{bmatrix} & + & 127 & 0 \\ Al_2O_3 \end{bmatrix} & + & Al_2O_3 \end{bmatrix}
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2[AlCl.] + 2[NaCl]
2[AlCl.] + 2[NH.[Cl]
2[AlCl.] + 3[KCl]
2[AlCl.] + 3[KCl]
2[AlCl.] + 6[KCl]
2[AlCl.] + 6[KCl]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            26 1 Cals.
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30 5
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            36.6
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5 3
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88 8
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Alloys.2 - Aluminium readily alloys with many other metals, and a large number of intermetallic compounds have been described. In general these compounds have little tendency to form continuous series of solid solutions with one another or with aluminum, and, accordingly, almost the only alloys of aluminium of engineering value are those which consist very largely of aluminium and those which contain but a few per cent of the element.3

Sodium and aluminium are not mutually soluble, beyond perhaps a very slight extent, and form no chemical compound.4

Copper and aluminium A number of investigations of the system copperaluminium have been made by the thermal method, but the results are somewhat contradictory. The compounds Al<sub>2</sub>Cu, AlCu, and AlCu<sub>3</sub> have been stated by several observers to exist. The "handus" curve has six or seven branches, only one of which appears to correspond to the separation of a pure compound, viz. Al2Cu. The other branches refer to the separation of solid

This number includes the least of solution of the anhydrous sulphate
 For a bibliography, complete to 1902, see Sack. Zedsch nanny Chem., 1903, 35, 249.
 For further information, see Law. Alloys (C. Griffin & Co., Ltd., 2nd ed., 1914), Gulliver, Alloys (C. Griffin & Co., Ltd., 2nd ed., 1913)
 Law. Trans. Faraday Soc., 1910, 6, 185
 Mathewson, Zedsch nanny. Chem., 1906, 48, 191.

solutions. 1 By a study of the electrical properties of these alloys, Broniewski has confirmed the existence of the compounds already mentioned, and in

addition has indicated the existence of another compound, Al, Cu,,

Aluminium copper alloys containing upwards of 10 per cent of aluminium have a fine golden-yellow colour, do not tarnish, are practically non-corrodible by sea-water, and are largely used under the name of aluminium bronzes. The tensile strength reaches the value 39 tons per sq. inch for the alloy with 10 per cent, of aluminium, the duetility is at a maximum with 7 per cent, of aluminium. The light copper-aluminium alloys containing upwards of 1 per cent. of copper are also of great value, by rolling and drawing, the tensile strength of the 4 per cent. alloy can be raised to 20 tons per sq inch (Carpenter and Edwards) A few per cent. of nickel are sometimes added to these alloys nich in aluminium, to obtain good rolling alloys

Silver and aluminium alloy containing 4 per cent. of silver has been used in the construction of balance beams; the alloy with 33 per cent has been employed for making tablespoons, etc. Two intermetallie compounds are known,  $Al\Lambda g_3$  and  $Al_2\Lambda g_3$ . They form a continuous series of solid solutions with one another, the former gives an incomplete series of solid solutions

with silver, and the latter an incomplete series with aluminium.8

Gold and aluminum Five intermetallic compounds are known, Au, Al,  $\mathbf{Au_8Al_3}$  or  $\mathbf{Au_2Al_2}$ ,  $\mathbf{Au_2Al}$ ,  $\mathbf{AuAl}$ , and  $\mathbf{AuAl_2}$ . The compounds  $\mathbf{Au_2Al}$  and  $\mathbf{AuAl_2}$  are stable at their melting-points, and the latter has a beautiful purple colour.4

Magnesium and aluminium. It is difficult to study this system by the thermal method. The "liquidus" appears to consist of three branches, the middle branch corresponding to the solid phase Al,Mg, or Al,Mg, middle branch, however, is extremely flat According to Broniewski, there are two intermetallic compounds, AlMg and AlMg, which form with one another a continuous series of solid solutions. The former gives an incomplete series of solid solutions with aluminium, and the latter an incomplete series with magnesium.5

Aluminium alloys containing 1-2 per cent of magnesium and a little copper, nickel, and tin are largely used for construction purposes under the name of magnatium, 6 the alloy duratumin contains 0 5 per cent. of magnesium, a few per cent of copper, iron, and manganese, and over 90 per cent. of aluminium.

<sup>&</sup>lt;sup>1</sup> Le Chatelier, Bull. Sw. d'Enc. p. Plud nationale, 1895, [iv.], 10, 509; Guillet, Compt. rend., 1901, 133, 684, 1914, 158, 701, Rev. Metallingie, 1905, 2, 567, Campbell and Mathews, J. Amer. Chem. Sw., 1902, 24, 253; Campbell, tind., 1904, 26, 1290, Carpenter and Edwards, Proc. Inst. Mech. Eng., 1907, p. 57, Curry, J. Physical Chem., 1907, 11, 425, Curry and Woods, ibid., 1907, 11, 461, Gwyei, Zeitsch anorg Chem., 1908, 57, 118, Portevin and Arnou, Compt. rend., 1912, 154, 511; J. H. Andrew, Intern. Zeitsch. Metallographic, 1414, 53.

and Arnou, Compt rend., 1912, 154, 511; J. H. Audrew, Intern. Zettsch. Mctallographie, 1914, 6, 30.

Bromewski, Ann. Chim. Phys., 1912, [viii.], 25. 5. This paper contains a useful summary of work on aluminum alloys, and also the results of investigations of the electrical properties of alloys of aluminum with numerous other metals.

1: Bromiewski, loc. cit.; cf. Petrenko, Zeitsch. anong Chem., 1905, 46, 49.

4: Heycock and Neville, Flid. Trans., 1900, A, 194, 201; 1914, A, 24, 267, Trans.

5Gem. Soc., 1898, 73, 714, Roberts-Austen, Proc. Roy. Soc., 1892, 50, 367; Matthey, 36dd., 1892, 51, 447, Osmond and Roberts-Austen, Proc. Roy. Soc., 1898, A, 187, 417.

5 Bromiewski, loc. cit., Grube, Zeitsch. anong Chem., 1905, 45, 225; Boudouard, Compt. rend., 1904, 138, 1501; Schimmester, Metallurgie, 1911, 8, 225, Pécheux, Compt. rend., 1904, 138, 1501; Schimmester, Metallurgie, 1911, 1, 522.

5 Eng. Pat., 1898, No. 24,878, Barnett, J. Soc. Chem. Ind., 1905, 24, 832, Klaudy, Oststr. Chem. Zeit., 1899, 2, 636.

Zinc and aluminium. It has been generally supposed that these metals form two series of solid solutions, the "liquidus" exhibiting a cutectic point, but Rosenhain and Archbutt have shown that there are three branches to the "liquidus," a compound Al<sub>2</sub>Zn<sub>3</sub> separating along the middle branch, which covers the region from 5-17 per cent of aluminium 1

The alloys known as riskon and risium are zinc-aluminium alloys, and are used for parts of scientific instruments. The alloys containing upwards of 20 per cent, of zine are valuable easting alloys, largely used in the motor industry. In actual practice, the amount of zinc seldom exceeds 10 per cent., and 2 per cent. of copper is added

The zinc-copper-alumnium alloys containing 31 27 per cent of zinc, 68-70 per cent. of copper, and 1-3 per cent of aluminium, form strong alloys, known commercially as aluminium brasses 2

Cadmium and aluminium, when fused and mixed, form two conjugate phases, one of aluminium containing 2 or 3 per cent of cadmium, the other of cadmium containing less than 1 per cent of aluminium 3

Mercury forms, with sufficient aluminium, a brittle solid amalgam which readily oxidises and decomposes water, forming mercury, aluminum hydroxide, and hydrogen. Aluminum is superficially amalgamated when it is immersed in mercuric chloride solution, and when so treated it rapidly decomposes water. The reaction is inhibited to a considerable extent by the presence of a little copper as impurity in the metal 6

Calcium and aluminium alloys have been studied, and the compound CaAl, isolated.6, 7

# ALUMINIUM AND THE FLUORINE GROUP.

Aluminium subfluoride, AlF2, is not known with certainty 8

Aluminium trifluoride, AlF, is not found in nature in the anhydrous state, but the hydrated fluoride occurs as the mineral fluellite, Alla, HaO. It crystallises in the rhombic system (bipyramidal , a:b:c=0.770:1:1:874); its density is 2 17

Aluminium is completely converted into the fluoride when heated in fluorine (Moissan) The fluoride is also formed when aluminium or alumina 10

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<sup>&</sup>lt;sup>1</sup> Rosenham and Archbutt, Phil Tians, 1911, A, 211, 315; Biomewski, loc cit.; Shepherd, J. Physical Chem., 1905, 9, 504; Pécheux Compt rend, 1904, 138, 1103; Smirnoff, that, 1912, 155, 551, Gautier, Bull Soc d'Em. p. Pind, nationale, 1896, [v.], 1, 1293; Heycock and Neville, Trans them Soc, 1897, 71, 383

<sup>2</sup> For an account of these alloys, see Carpenter and Edwards, Intern Zeitsch Metallographic, 1912, 2, 209, Levi-Malvano and Mirantonio, Gazzita, 1911, 41, 1–282, 1912, 42, i. 353

For aluminium-zine-magnesium alloys, see Egei, Intern Zeitsch Metallographic, 1913 a 29

<sup>1913, 4, 29.

\*</sup> Wright, J. Soc. Chem. Ind., 1892, II. 192, 1894, I3, 1014, Campbell and Mathows,
J. Amer. Chem. Soc., 1902, 24, 253

\* Baille and Féry, Ann. Chem. Phys., 1889, [v1], 17, 246, Biernacki, Wied. Annalen,
1896, 59, 664, Humphreys, Trans. Chem. Soc., 1896, 69, 1679

\* Kohn-Abrest, Indl. Soc., chim., 1910, [v], 7, 283, 1912, [v], II, 570, Kohn-Abrest,
and Rivota-Maltes, Compl., rend., 1912, 154, 1600, Nicolardot, Bull. Soc., chim., 1912,
[iv.], II, 410; see also p. 379.

\* Donski, Zeitsch. anorg Chem., 1908, 57, 185

\* Gullium aluminium alloys are mentioned on p. 147, thallium-aluminium alloys on p. 175; ceriam-aluminium alloys, 3, 1; Blackmore, J. Soc. Chem. Ind., 1897, 16, 219

\* Poulene, Ann. Chim. Phys., 1894, [vii.], II, 66.

Brunner, Pogg. Annalen, 1856, 98, 488, van Haagen & Smith, J. Amer. Chem. Soc.,
1911, 33, 1504.

is heated to redness in hydrogen fluoride. Deville, who made an exhaustive study of the fluoride, obtained it first by heating aluminium in silicon fluoride, and afterwards prepared it by treating alumina with hydrofluoric acid and subliming the product in hydrogen at a white heat. He also obtained the fluoride by heating a mixture of fluorspar and alumina in a stream of hydrogen fluoride, and by fusing cryolite with anhydrous aluminium sulphate.1 The impure fluoride is best purified by sublimation at 1100°.

Anhydrous alummum fluoride forms colourless, transparent crystals which are described as rhombohedra,2 but, according to Poulenc, are probably trichme. It is exceedingly refractory, being insoluble in water and unattacked by alkalies and acids, including concentrated sulphuric acid. It may

be decomposed by prolonged fusion with an alkali carbonate.

When 42 grams of basic aluminium acetate, 51 grams of alumina, 100 grams of water, and 66 grams of 40 per cent hydrofluoric acid are mixed and warmed, complete solution results, and a gelatmous, hydrated aluminium fluoride separates on standing If the solution be somewhat diluted, crystals of the trihydrate, AlF, 311,0, slowly separate, if only 75 grams of water are employed in the preparation, crystals of another hydrate, 2AH, 17H,0, separate. The latter hydrate effloresces in air, passing into the former. The monohydrate, AIF, II\_O, can also be obtained in spannigly soluble, silky needles, and likewise the hydrate, 2AIF, 7II\_O, which slowly changes to the

hexahydrate, AH<sub>3</sub> 6H<sub>4</sub>O, on standing in water <sup>3</sup>
The hydrate 2 MF<sub>4</sub>7H<sub>4</sub>O has been obtained by Baud in two forms first, practically insoluble in water, is obtained by dissolving aluminium hydroxide in aqueous hydrofluoric acid and evaporating the solution at 100°, the second, easily soluble in water, by concentrating a solution of aluminium hydroxide in hydrofluoric acid and adding twice its volume of alcohol. An The hydrate 2 MF , 7H2O aqueous solution of the latter form is acid to litmus loses water at 110°, leaving the hemitydrate 2AIF, If,O, which decomposes

at a bright red heat's

According to Deville, the compounds AF, 3HF, 3AF, 2HF 5H,0, and 2AF, HF.5H,0 may be prepared. Aluminium fluoride enters into the composition of a large number of double compounds. With each of the alkali fluorides it forms a compound of the type AIF, 3NF stable at its melting- ${\bf point.}$  The melting points are given by Puschin and Baskov  $^5$  as follows -

Alkali Metal Rb Na Melting-point of Salt, °C . 800° 1035 1020° 985°

According to the same authors, compounds of the type 2AlF, 3NF are also formed when X -- Na, K, or Rb, but this conclusion is not justified from their experiments and is probably erroneous. In the case of the sodium aluminium fluorides, the second compound has the formula 3AIF, 5NaF, and has no melting-point, but dissociates at 723° into AIF3.3NaF and aluminium fluoride.

Deville, Compt. rend., 1856, 42, 19., Ann. Chim. Phys., 1857, [ii ], 49, 79, 1861, [iii], 61, 333. For other methods, see Hautefeuille, ibid., 1805, [iv ], 4, 153. Troost and Hautefeuille, Compt. rend., 1872, 75, 1819, Friedel, Bull. Soc. chim., 1874, 21, 241; Cossa, Gazzetta, 1877, 7, 212.
 De Schulten, Compt. rend., 1911, 152, 1261
 Mazzucchelli, Atts R. Accad. Lincet, 1907, [v ], 16, 1, 775.
 Baud, Ann. Chim. Phys., 1904, [viii ], 1, 8
 Puschin and Baskov, J. Russ. Phys. Chem. Soc., 1913, 45, 82; Zeitsch. anorg. Chem., 1913, 87, 347.

<sup>•</sup> Fedotéev and Iljinsky, Zeitsch. anorg. Chem ; 1913, 80, 113.

The equilibrium diagram, as far as it has been worked out, is shown in fig. 8. Calcium and aluminium fluorides form no compound; the eutectic point

Hydrated double fluorides of the formulæ 2AlF<sub>8</sub> 6KF.7H<sub>2</sub>O, 2AlF<sub>8</sub>6NaF. 7H2O, and 2AlF3.4NH4F.3H2O are obtained as gelatinous precipitates when the requisite alkali fluoride solutions are added to an aqueous solution of Baud's soluble hydrate of alummium fluoride They are slightly soluble in water, 100 parts of which dissolve 0.385 of the potassium, 0.352 of the sodium, and 1 0 of the ammonium compound at 16° (Baud, loc. cit.) Berzelius has described an insoluble double fluoride AlF, 3NH,F, and Petersen and Helmolt have prepared a soluble double fluoride of the same composition.

The double fluoride 3AIF, 5NaF occurs in nature as the tetragonal mmeral chiolite, the compound AIF, 3NaF is found as the mineral cryolste. Cryolite ( & e. 1ce-stone) occurs at lyigtut, an Esquimaux hamlet on the south west coast of Greenland, in one huge deposit, contaminated with siderite, zincblende, galena, etc. Density, 2.96, hardness, 2.5, melting-point, c. 1000°. It forms monoclinic prisms (holohedral, a b c = 0.9662·1:13882,  $\beta$  = 90° 11') 2 and at c .565° is converted into a cubic modification. At 15°, 100 parts of water dissolve 0 034 of cryolite (Baud), but the latter is appreciably soluble in aqueous solutions of aluminium salts It is decomposed by sulphuric acid. In the purification of crude cryolite, advantage is taken of the superior density of its impurities to effect a first punification, and other impurities are then removed by means of an electromagnet. Cryolite is used as a solvent for alumina in the process of manufacturing aluminium (p. 48), in the preparation of opaque white glass, and of an enamel for steel. It was formerly used as raw material for the Danish alkalı industry (see Vol II.), and ha been powdered and used by the Esquimaux as snuff 3

The following double salts may be prepared by dissolving the requisite hydroxides in hydrofluoric acid and evaporating the solution -

 $\begin{array}{c} {\rm AlF_4~ZnF_2.7H_2O} \\ {\rm AlF_3.2CuF_2.11H_2O} \end{array}$  $\substack{2\text{AIF}_3\text{ 3CuF}_2\text{ 18H}_2\text{O}\\\text{AIF}_3\text{ CuF}_2\text{ IIF }8\text{H}_2\text{O}}$ 

Aluminium subchloride has been said to be produced by heating aluminium trichloride with aluminium in a scaled tube, but the statement is in all probability erroneous.6

Aluminium trichloride, AlCI, was originally made by heating an intimate mixture of alumina and carbon to iedness in a stream of chlorine (Oersted's method)? It may be more readily prepared by heating aluminium in a wide glass tube in a rapid current of dry hydrogen chloride, or in a stream of chlorine 8 If it is required to prepare the chloride from the oxide, a

<sup>1</sup> Petersen, J prakt Chem., 1889, [11], 40, 55; Helmolt, Zeitsch. anorg. Chem., 1893, 3, 115. On the detection of sodium by piccepitation as sodium aluminum fluoride, see Wilks, Proc. Camb. Phil. Soc., 1909, 15, 76

2 Krenner, Zeitsch. Kryst. Min., 1885, 10, 525.

3 See Halland, J. Ind. Eng. Chem., 1911, 3, 63.

4 Weinland and Koppen, Zeitsch. anorg. Chem., 1899, 22, 266.

5 Friedel and Roux, Compt. rend., 1885, 100, 1191.

6 Nilson and Petterson, Trans Chem. Soc., 1888, 53, 826

7 Wohler, Poyg. Annalen, 1827, 11, 146. Leebig, ibid., 1830, 18, 43; Bursen, ibid., 1854, 92, 648. Deville, Compt. rend., 1849, 29, 321, Ann. Chim. Phys., 1855, [in.], 43, 11.

8 Stockhausen and Gattermann, Ber., 1892, 25, 3521, Nilson and Petterson, vide. 1847; Escales, Ber., 1897, 30, 1314, Gustavson, J. prakt. Chem., 1901, [in.], 63, 110; Köhn-Abrest, Bull. Soc. chim., 1909, [iv.], 768.

neater method than Oersted's is to heat the oxide in a current of chlorine and sulphur chloride :- 1

$$4\text{Al}_2\text{O}_3 + 3\text{S}_2\text{Cl}_2 + 9\text{Cl}_2 = 8\text{AlCl}_3 + 6\text{SO}_2$$

Instead of chlorine and sulphur chloride, carbon tetrachloride vapour  $^2$  or carbonyl chloride  $^3$  may be used. A simple method of preparation is said to consist in heating crude alumina or clay to redness in a current of hydrogen chloride and carbon disulphide vapour, and punifying the aluminium chloride so obtained by sublimation over non filings,4

Aluminium chloride, purified by sublimation over aluminium, forms white, lustrous, six-sided plates which are said by Seubert and Pollard to possess rhombic symmetry The slightly impure chloride is usually yellow owing to the presence of a little ferric chloride When slowly heated, aluminium chloride sublimes completely, but when a mass of the chloride is rapidly heated it molts.

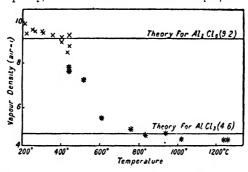


Fig. 11.—Vapour density of aluminium chloride – Results by Dumas' method shown  $\times$  , results by Victor Meyer's method shown  $^{\bullet}$ 

The sublimation pressure of the chloride reaches one atmosphere at 179° to 183°; the melting point at 2.5 atmospheres pressure is  $190^{\circ}$  to  $194^{\circ}$ . The critical temperature is  $629.5^{\circ}$  C.6°. The vapour density of aluminium chloride has been the subject of numerous researches. The results that have been obtained at atmospheric pressure and different temperatures (method of Dumas) are shown by crosses in fig. 11. The asterisks in the same figure

- 1 Matignon and Bonton, Compt. rend., 1901, 138, 631, 760, Bourton, Ann. Chim. Phys., 1910, [vnt], 20, 547. Alumina is not attacked by SOCI, at 200° (North and Hageman, J. Amer. Chem. Soc., 1913, 35, 352)

  2 Demarcay, Compt. rend., 1837, 104, 111, Quantin, ibid., 1888, 106, 1074; Camboulives, ibid., 1910, 150, 175, 221; L. Meyer, Ber., 1887, 20, 681.

  3 Chauvenet, Compt., cond., 1911, 152, 87

  4 Curic, Chem. News, 1873, 28, 307. For other methods of preparation, see Rose, Pogy Annales, 1848, 74, 569, Havitsky, Ber., 1873, 6, 195, Thost and Hautelle, Compt. rend., 1872, 75, 1710, 1819; Weber, Pogy Annales, 1857, 101, 165, 165, 103, 259, Warren, Chem. News, 1887, 55, 192; Faure, Compt. rend., 1888, 107, 339, Mabery, Ber., 1888, 22, 2058.

  5 Friedel and Crafts, Compt. rend., 1888, 106, 1764; Seubert and Pollaul, Ber., 1891, 24, 2575.

- Friedel and Cratts, Compt. Tena., 1000, 100, 1101, 1014, 87, 635
   Rotmanz and Suchodski, Zeitsch. physikal. Chem., 1914, 87, 635
   Deville and Troost, Ann Chim Phys., 1860, [in.], 58, 257; Nilson and Pettersson, Officers. K Stenska Vet. Akad. Firhandl., 1887, No. 8, Zeitsch physikal. Chem., 1887, 1, 459; Ann Chim. Phys., 1890, [vi.], 19, 145, Friedel and Grafts, loc. cit., Friedel, Ann. Chim. Phys., 1890, [vi.], 19, 171.

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represent the values obtained by the method of Victor Meyer, in which the partial pressure of the aluminium chloride was less than one atmosphere, but had no definite value throughout a series of experiments. The theoretical value for  $Al_a Cl_b$  is 9.2 (air = 1), and for  $Al Cl_b$  4.6. It will be seen that from 200° to 400° the molecular formula is  $Al_a Cl_b$ , above 400° the dissociation  $Al_a Cl_b == 2A|Cl_b$  becomes marked, and at and above 800° the dissociation is complete. The specific heat of aluminium chloride is 0 188.1

Alummium chloride dissolves in alcohol and many organic liquids. In pyridine, ether, and in introbenzene the molecular weight corresponds with the simple formula AlCla 2

Alummum chloride is extremely deliquescent and fumes in the air. It dissolves readily in water with the evolution of much heat (p. 61). An aqueous solution of aluminium chloride is readily obtained by dissolving aluminium or its hydroxide in aqueous hydrochloric acid. A solution containing 40 parts of aluminium chloride to 100 of water has a density of 1.3415 at  $15^{\circ}$ . The aqueous solution has an acid reaction and on prolonged boiling loses hydrochloric acid and becomes turbid. According to Baud, a number of definite basic chlorides exist !

The hexahydrate, AlCl<sub>3</sub>.6H<sub>2</sub>O, separates out when an aqueous solution of aluminium chloride is either slowly evaporated or saturated with hydrogen chloride The hydrate forms deliquescent rhombohedral crystals (a c=1 05356) of prismatic habit and does not lose water in a sulphuric acid desiccator. Dilute solutions, cooled to  $-8^{\circ}$ , are said to deposit the ennea-hydrate AlCl, 9H<sub>2</sub>O.7

Aluminium chloride enters into the composition of a large number of double compounds, many of which have been studied by Band 8. The following compounds with metallic chlorides are known -

```
2AlCl<sub>F</sub>6KCl
2AlCl<sub>F</sub>2AgCl
2AlCl<sub>F</sub>BaCl<sub>2</sub>
2AlCl<sub>F</sub>1½SrCl<sub>2</sub>
2AlCl<sub>F</sub>1½CaCl<sub>2</sub>
2AlCl, 2NH<sub>1</sub>Cl
2AICI, 2KCI
2AICI, 2NaCI
2AICI, 3NaCI
2AICI, 3KCI
2AICI, 6NaCl
  2AlCl, 1½ZnCl2
```

The compound AlCl NaCl was formerly manufactured for use in the preparation of aluminum. It is not so hygioscopic as aluminum chloride, melts at 185, and volatilises at a red heat. The compounds  $\Lambda ICI_3$  KCl and AlCl, NII, Cl are similar

The compound AlCl3 SCI4 has been prepared by Ruff and Plato, double compounds with sclemin and tellurium tetrachlorides by Weber, and with phosphorus pentachloride and oxychloride by Casselmann.

<sup>1</sup> Baud, Ann Chim Phys, 1994, [viii.], I, 8.
2 Werner and Schmuplow, Ziisch anorg Chem., 1897, I5, 24, Beckmann, Zeitsch.
physikal. Chem., 1903, 46, 860, Kohler, Imer Chem. J., 1900, 24, 335, respectively.
3 Gerlach, Zeitsch. onal. Chim., 1869, 8, 250
4 Baud, loc. cit. See also Lacehit and Supla, J. Nor Chim. Ind., 1883, 2, 539, Tommasi,
Bull. Soc. chim., 1882, 37, 443, Hautefeuille and Perrey, Compt. rend., 1885, 100, 1219;
Schlumberger, Bull. Soc. chim., 1895, [iii], 13, 48
5 Bonsdouff, Porg. Annalen, 1833, 27, 279
6 Dennis and Gill, Zeitsch. anorg Chem., 1895, 9, 339.
7 Lubaiski, Zeitsch. anorg. Chem., 1898, 18, 387
8 Baud, Ann. Chim. Phys., 1901, [viii], I, 8, see also p. 61
9 Ruff and Plato, Ber., 1901, 34, 1749; Weber, Porgs. Annalen, 1858, 104, 421;
Casselmann, Annalen 1856, 98, 220.

Anhydrous aluminium chloride rapidly absorbs dry ammonia. At low temperatures the compound AlCl<sub>3</sub> 9NH<sub>3</sub> is formed, the dissociation-point of which is -146°. It passes into the compound AlCl<sub>3</sub>.6NH<sub>3</sub>, which is the product formed at the ordinary temperature and pressure At 150° this passes into AlCl<sub>3</sub> 5NH<sub>3</sub>; at 275° this becomes AlCl<sub>3</sub> 3NH<sub>3</sub>; and when this last compound is distilled in hydrogen, a compound of the remarkable composition 6AlCl<sub>3</sub>·7NH<sub>3</sub> is produced which may be repeatedly distilled in hydrogen without decomposition <sup>1</sup> The compounds AlCl<sub>3</sub>·XCl 6NH<sub>3</sub> (where X = NH<sub>4</sub>, Na, or K) are also known <sup>1</sup> Aluminium chloride also combines with phosphine.

The compound  $AlCl_3$   $II_2S$  can only be obtained by the use of liquid hydrogen sulphide  $\,$  It dissociates at  $-45^\circ$  into hydrogen sulphide and the compound 2AlCls. H2S, which is stable at the ordinary temperature and pressure The compound AlCl<sub>3</sub> SO<sub>2</sub>, prepared by subliming aluminium chloride in a current of sulphur dioxide, dissociates at 80°, giving rise to the compound 2AlCl<sub>3</sub> SO<sub>2</sub>, which can be distilled at 200°.2

With carbonyl chloride three compounds are formed, 2AlCl<sub>3</sub>.5COCl<sub>2</sub>, melting at -2°, 2AlCl, 3COCl, and 4AlCl, COCl. The last two compounds are said to occur in commercial aluminium chloride.8

The behaviour of the preceding double compounds towards water is such as would be expected from the behaviour of the constituent compounds.

Aluminium chloride combines with many organic compounds, such as acid chlorides, ketones, estons, intro-compounds, and tertiary amines. Examples of such compounds are AlCl<sub>3</sub>.(C<sub>2</sub>H<sub>5</sub>),O, AlCl<sub>3</sub> C<sub>6</sub>H<sub>5</sub>,CO<sub>2</sub>C<sub>2</sub>H<sub>5</sub>, AlCl<sub>3</sub>.NO<sub>2</sub> C<sub>6</sub>H<sub>4</sub> CH<sub>3</sub>, Al<sub>2</sub>Cl<sub>6</sub> 2C<sub>6</sub>ll<sub>5</sub>NO<sub>2</sub>, the last compound having a molecular weight in carbon disulphide solution corresponding to the formula here given.4 In organic chemistry, anhydrous aluminium chloride is a very valuable catalytic agent, as, for example, in the well-known Friedel and Crafts' syntheses.5

Aluminium tribromide, AlBr,, is most readily prepared by adding aluminium carefully, in small quantities at a time, to bromine, the reaction being extremely vigorous 6 Other methods of preparation are to pass bromme vapour over heated alummium or an intimate mixture of alumina and carbon heated to redness.7 The product should be rectified over metallic aluminium.

Aluminium tribromide forms colourless, transparent rhomboliedia of density 2.54. It melts at 93° and boils at 263°3° under 747 inm. pressure (Mallet), its critical temperature is 772° C.8 At 444° and 1 atmosphere the vapour density is 1862, the formula Al<sub>2</sub>Br<sub>6</sub> corresponding to the value 18 12 9 The specific heat of aluminium brounde (22" to 76") is 0.08912, and the latent heat of fusion 10.47 cals. per gram , as a cryoscopic solvent the molecular depression of the freezing-point is  $262^{-10}$  The bromide is a non-electrolyte, but dissolves numerous halogen salts with the production of conducting solutions.11 Aluminium bromide dissolves in alcohol

Baud, loc. cit; cf. Stillman and Yoder, Amer. Chem. J., 1895, 17, 748.

<sup>Baud, loc. cit.
Baud, Compt. rend., 1905, 140, 1688.
Kohler, Ames. Chem. J., 1900, 24, 385.
Friedel and Crafts, Compt. rend., 1877, 84, 1392, 1150, 85, 74, 672.
Mallet, Phil. Trans., 1880, 171, 1003
Deville and Troost, Ann Chim. Phys., 1860, [iii.], 58, 257; Gustavson, J. prakt. Chem., 1901, [ii.], 63, 110; Kablukoff, J. Russ. Phys. Chem. Noc., 1908, 40, 485.
Rotnjanz and Suchodski, Zeitsch. physikal. Chem., 1914, 87, 635.
Daville and Troost. Loc. cit</sup> 

Rotinjanz and translation
 Daville and Troost, loc cit
 Kablukoff, loc cit.; Isbekoff, Zeitsch. anorg. Chem., 1913, 84, 24.
 Kablukoff, J. Russ. Phys. Chen. Soc., 1911, 43, 18.

and many organic liquids The molecular weight corresponds to the double formula Al2Br, in carbon disulphide 1 and in biomine, 2 but in nitrobenzene it corresponds to the single formula AlBr, 1 From a solution of aluminium bromide in ethyl bromide, aluminium can be deposited by electrolysis, but the deposited metal readily attacks the solvent, butane being produced 3

Aluminum brounds is decomposed when heated in oxygen. It is extremely deliquescent and dissolves in water with the evolution of much heat. The aqueous solution, which may be prepared from alumnium or its hydrovide and hydrobromic acid, reacts acid and becomes basic when boiled. The hexahydrate AlBr3.6H2O can be prepared by methods corresponding to those given for the hevahydrate of aluminium chloride (p. 67); a dilute aqueous solution when cooled is said to deposit the hydrate 2AlBr, 15H2O.4

Aluminium bromide forms double salts with the alkali bromides, of which AlBr, KBr and AlBr, NaBr have been described. With the tribroundes of antimony and bismuth it forms solid solutions 6 It combines with hydrogen sulphide, forming a white, crystalline compound, AlBr, II S, which melts at 84°, 7 and with phosphorus ovychloride, forming a compound of the molecular formula  $Al_2Br_0.2PO(l_1)$  in carbon disulphide solution, The compounds  $AlBr_1.2Br_2.CS_2$  and  $2AlBr_1.2Br_2.CS_2$  have also been described. Like the chloride, it combines with many organic compounds The following are the molecular formulæ, in carbon disulphide solution, of a few such compounds:-1 ${\rm Al_2Br_6.2C_6H_5SO_2Cl,\ Al_2Br_6.2(C_6H_5)_2CO,\ Al_2Br_6.2C_6H_5COCl,\ Al_2Br_6\ 2C_6H_5NO_2.}$ 

Aluminium iodide, All,, may be prepared by heating aluminium with iodino in a scaled tube, by passing iodino vapour over heated aluminium, or by adding aluminium to iodine dissolved in carbon disulphide. It separates from carbon disulphide in colourless crystals of density 2.63, melts at 125°, and boils at 350°, its critical temperature is 955° 10° At 111° the vapour density is 27 0 (air = 1), the formula Algla requiring 28 2 (Deville and Troost). The vapour forms an exclusive mixture with air. The molecular formula is Al2I6 in molten iodine,11 and in carbon disulphide solution.

Aluminum iodulo is very soluble in liquid animonia, from which the compound AlI<sub>3</sub>.20NII<sub>3</sub> (?) may be crystallised at -33° (° At 8-13° C, the crystals lose ammonia and leave the compound AlI<sub>3</sub> 6NII<sub>3</sub>. The liquid annuonia solution reacts with potassium annuc to form a soluble aluminium ammonobasic iodide, Al(NH2), All, which crystallises with six molecules of ammonia at the ordinary temperature and about twenty at low temperatures. This compound reacts with more potassium amide to produce an insoluble ammonobasic rodide, Al(NH2), Al(NH2), I.NH, which loses two molecules of ammonia at 160° C 12

Kohler, Amer Chem. J., 1900, 24, 385.
 Beckmann, Zeitsch. anory. Chem., 1906, 51, 96.
 Plotnikoff, J. Russ. Phys. Chem. Soc., 1902, 34, 466, Patten, J. Physical Chem., 9, 2849. 1904, 8, 548.

<sup>Weber, loc. cet; Deville and Troost, Ann. Chim. Phys., 1860, [iii], 58, 257;
Gustavson, Annalen, 1874, 172, 173.
Rotinjanz and Suchodski, Zeitsch physikal Chem., 1914, 87, 635.
Beckmann, Zeitsch. anorg. Chem., 1912, 77, 200, 275.
Franklin, J. Amer. Chem. Soc., 1915, 37, 847.</sup> 

Aluminium iodide is very soluble in water, an aqueous solution can be readily prepared from aluminium or its hydroxide and hydrodic acid the chloride and bromide, it forms a hexahydrate, All, 611,0, it also forms a hydrate, 2AH, 15H,O1

The double salts All, KI, All, NaI and All, HgI, 8H,O are known.2

Aluminium chlorate, Al(ClO<sub>1</sub>), 911,O, prepared by mixing aluminium sulphate and barum chlorate, removing the barum sulphate, and evaporating the cold solution over sulphune and, is very soluble in water. From a hot solution it separates as the hexahydrate. The salt is easily decomposed by heat, and explodes when slowly heated.3

Aluminium perchlorate, Al(ClO<sub>1</sub>), 6H<sub>2</sub>O, is a colourless, crystalline salt, soluble in water. Its aqueous solution gives, with sodium perchlorate, a crystalline precipitate of aluminium sodium perchlorate, AlNa(ClO<sub>1</sub>), 12H<sub>2</sub>O.4

Aluminium bromate, Al(BrO<sub>3</sub>), 911.0, is prepared like the chlorate. It melts at 62.3° and easily decomposes 5

Aluminium periodate, Al(101), 31120, crystallises in regular octahedra, 6

### ALUMINIUM AND THE OXYGEN GROUP.

Aluminium suboxide. - The existence of a suboxide of aluminium has been suspected by several chemists 7

Aluminium sesqui-oxide or alumina, Al<sub>2</sub>O<sub>3</sub>, is found in the crystalline form in nature as the mineral conundum. It crystallises in the trigonal system (ditrigonal scalenohedral;  $a=c=1-1.365)^8$  usually in double, six-sided pyramids and thombohedra, with the basal plane. It is therefore isomorphous with ferric and chromic oxides Haidness, 9, density, 3.9-41; lustre, viticous. The dull and opaque varieties, or "common corundum," occur in India, China, Siberia, and the United States, and are largely used as abrading agents. The transparent varieties are highly valued as gem-stones. Clear, colourless stones are known as white suppliere; blue stones as sapphere, red stones as oriental ruby, 10 yellow stones as oriental topaz or yellow sapphire , purple stones as oriental amethyst , and the rare, green stones as oriental emerald. These coloured varieties of corundum are pleochroic. Sapphires are found in Ceylon, Burma, Stain, and parts of India, and in the gold-bearing drifts of Victoria and New South Wales. Oriental rubics occur in Ceylon and also at Mogok, Upper Burma 11

Impure granular or crystalline corundum, associated with magnetite, tourmaline, gainet, etc., occurs in nature as emery. It is obtained from

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1 Panfiloff, J. Russ. Phys. Chem. Soc., 1895, 27, 77.
2 Weber, toe cit., Dubou, Compt. rend., 1908, 146, 1027.
3 Dobroserdoff, J. Russ. Phys. Chem. Soc., 1904, 36, 468
4 Weinland and Ensgraber, Zeit-ch. anorg. Chem., 1913, 84, 368.
5 Dobroserdoff, J. Russ. Phys. Chem. Soc., 1907, 39, 133
6 Eakle, Zeitsch Kryst. Min., 1890, 26, 558.
7 Pronchon, Compt. rend., 1893, 117, 328, Kohn-Abrest, Bull. Soc. chim., 1901, [in], 31, 323; Compt. rend., 1905, 144, 323, Duboin, ibid., 1901, 132, 826.
6 Melcer, Zeitsch Kryst. Min., 1902, 35, 561.
9 Jenks, Quart. J. Geol. Soc., 1874, 30, 303, Judd and Hidden, Min. Mag., 1899, 12, 139; J. H. Pratt, Bull. US. Geol. Survey, 1906, No. 269
10 Various other real stones are technically termed ruby, eq. spinel, garnet, hence the qualification "oriental" applied to corundum rubies.
11 C. W. Brown and Judd, Phil. Trans., 1896@A, 187, 151; Judd, Min. Mag., 1895, 11, 56; Louis, ibid., 1894, 10, 267.
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71 ALUMINIUM.

Navos in the Greek Archipelago, Asia Minor, and Massachusetts, and is used as an abrading agent,

Alumina is obtained as an amorphous white powder or guin-like mass by heating aluminum hydrovide or the aluminium salt of a volatile oxyacid, e.g. the mitrate, sulphate, ctc When molten alumina solidifies, when alumina vapour condenses to the solid state and when alimina separates from its solution in a suitable solvent, it assumes the crystalline form and characteristics of coundum. Rhombohedral crystals may be obtained by heating amorphous alumma with five times its weight of aluminium sulphide in the elective furnace, and treating the product with hydrochloric acid 2

The crystallisation of alumina has been attempted by various chemists in the hope of preparing rubies and sapplines. The ruby owes its colour to a trace of chromic oxide. The first successful experiments on the production of rubies on a large scale were made by Frémy and Feil,3 who fused equal parts of alumina and bitharge, plus 2 or 3 per cent of potassium dichromate, in a fireday crucible at a bright red heat. The product consisted of a layer of lead silicate and a viticous layer in which crystals of ruby were embedded. Later, by replacing the lithinge by barrum fluoride and heating in a glass furnace, Fremy and Verneurl obtained beautiful rubics, which had arisen by the action of the furnace gases on altanuum fluoride vapour! Loyer obtained rubies by heating sodium aluminate (100 pts.) and potassium dichromate (1 pt.) to bright redness in chlorino.

At the present time, rubies are manufactured by a process devised by Vernoul <sup>6</sup>. The material used is powdered alimina containing a little chromo oxide (2.5 per cent ), produced by precipitating with ammonia a solution of pure ammonium alum to which a little chrome alum has been added, and igniting the mixed hydrovides. The powder is fed through the oxygen tube of an inverted oxy-coal-gas blowpipe, falls as a molten drop on to the end of a small alumina rod, and crystallises as ruby—As the process is continued, the ruby grows upwards as a pear shaped drop or "boule"—These "boules" have a density of 401, although externally smooth, they have a crystalline structure and differ from natural stones only in one respect, viz, they contam microscopic air-bubbles and fine-curved internal streaks. The streaks in a natural stone are straight?

Sapphires are manufactured from alumna to which 15 per cent. of magnetic non oxide and 0.5 per cent of titamium oxide has been added, the mixture being fused and crystallised in an oxy-hydrogen reducing flame 8 The introduction of cobalt oxide into alumina can only be effected in the presence of a third oxide such as lime. The stones obtained are blue, but

<sup>1</sup> A second form of alumna is sometimes obtained when molten alumna is slowly cooled,

<sup>&</sup>lt;sup>2</sup> A second form of alumina is sometimes obtained when molten alumina is slowly cooled, particularly in the presence of a little magnesia. It is hexagonal, density 3.80. See Rankin and Mewin, J. Lines. Chem. Sov. 1916, 38, 570.

<sup>2</sup> Hondard, Compt. rend., 1907, 144, 1319

<sup>3</sup> Frémy and Penl. Compt. rend., 1877, 85, 1029; Thil. May., 1878, [v.], 5, 47.

<sup>4</sup> Frémy and Venneul, Compt. rend., 1887, 104, 737, 738, 1888, 106, 565, 1890, 111, 667, Desclorzeaux, thid., 1888, 106, 567.

<sup>6</sup> Loyer, Bull. Sov. chim., 1897, [int.], 17, 345

<sup>6</sup> Venneul, Ann. Chem. Phys., 1904, [vim.], 3, 20.

<sup>7</sup> For early work on the ruby, see Gaudin, Compt. cend., 1837, 4, 999; 1857, 44, 716; 1869, 69, 1313, Ebelmen, Ann. Chim. Phys., 1851, [in.], 33, 34, Deville and Caron, viola, 1865, [iv.], 5, 104. Hautefeulle, thid. 1865, [iv.], 4, 153., Debray, Compt. cend., 1861, 22, 985; De Semamont, thid., 1851, 32, 762., Deville, thid., 1855, 40, 771. See also Friedel, Bull. Soc. chim., 1886, [in.], 46, 242.

<sup>8</sup> Vernentl, Compt. rend., 1910, 1504, 185.

#### ALUMINIUM AND ITS CONGENERS.

A ... A . A ...

they are amorphous, and have not the tint of sapphire.1 The analyses of three natural sapphires gave the following results -

| Locality      | Australia, | India. | Montana |
|---------------|------------|--------|---------|
| Fe,(),        | 0.920      | 0.72   | 0.560   |
| TıÒ, "        | 0 031      | 0.04   | 0.058   |
| SiO,          | trace      | nıl    | 0 100   |
| Al,Ö, (dıff.) | •••        |        |         |

It has been shown that synthetic and natural sapphires have identical properties.3

About ten million carats of rubies and six million carats of sapphires were manufactured in 1913, and the demand is increasing.4

Amorphous alumina is a white powder, insoluble in water. The density increases with the temperature at which it has been ignited, as follows .-- 5

| Temperature ° C | . 600° | 700° | 800° | 900° | 1200° |
|-----------------|--------|------|------|------|-------|
| Density         | 282    | 2 83 | 3 39 | 3 53 | 392   |

When slowly heated it undergoes an exothermic change at 850°,6 and inelts at 2010° to 2050.7 In the electric furnace it melts and boils, and the vapour condenses to the crystalline form.8 The specific heat of ignited alumina increases with the temperature, as is shown by the following mean values -

Alumina that has been dried at a low temperature is very hygroscopic and forms an excellent drying agent.10

Alumina is soluble in mineral acids unless it has been strongly heated (above 850), when it becomes extremely refractory. Calcined alumina must be brought into solution by fusion with potassium hydrogen sulphate or alkali hydroxide.

Alumina is unaffected by hydrogen or chlorine at a red heat, but is converted by fluorine into aluminium fluoride and ovygen. At a red heat it is converted into the sulphide by carbon disulphide. Alumina is reduced by carbon to the metal at temperatures above 2000', 11 i.e. at temperatures above which aluminium carbide is unstable 12 It was shown by Moissan that carbon reduces alumma vapour.13

Alumina is used in the manufacture of aluminium, for this purpose it is prepared by igniting the hydroxide, prepared from banxite as described later

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    Paris, Compt. rend., 1908, 147, 933; Verneuil, 1bid., 1908, 147, 1059.
    Verneuil, Compt. rend., 1910, 151, 1063.
    Moses, Amer. J. Sci., 1910, [iv], 30, 271.
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<sup>\*</sup> For a review of the subject of synthetic stones, see Levin, J. Ind. Eng. Chem., 1913,

<sup>\*\*</sup> For a leview of the subject of synthetic stories, 5.

5. 495.

\*\* Mellor and Holderoft, Trans. Eng. Ceramic. Soc., 1911, 9, 94.

\*\* Le Chatelier, Compt. vend., 1887, 104, 1517; Bull. Noc. chim., 1887, [iii], 47, 300.

\*\* Kanolt, J. Washington Acad. Sci., 1912, 2, 337., 1913, 3, 315; Ruif and Goecke, Edisch. angew. Chem., 1911, 24, 1459; Ruif, Serferheld, and Suda, Zeitsch. anorg Chem., 1918, 82, 873; Tiede and Birubrauer, ibid., 1914, 87, 129.

\*\* Moissan, Compt. rend., 1892, 115, 1034.

\*\* Tilden, Trans. Chem. Soc., 1905, 87, 551.

10 Johnson, J. Amer. Chem. Soc., 1912, 34, 911.

11 Hutton and Petavel, Proc. Roy Soc., 1907, A, 79, 155.

12 Askensay and Lebedeff, Zeitsch. Elektrochem., 1910, 16, 559.

13 Moissan, Compt. rend., 1894, 119, 935; Bull. Soc chim., 1895, [iii], 13, 803.

(p. 79). It has been proposed to prepare it from aluminium nitride, manufactured by Serpek's method (p. 88). Another proposal, which may possibly develop into a successful commercial method, is to obtain the alumina from sodium aluminate, itself manufactured from china-clay.1

The porous alundum laboratory utensils are composed mainly of alumina. Calcined bauxite is fused in a water-cooled electric are furnace. The impurities in the bauxite are to a certain extent reduced and segregate at the bottom of the fused mass as an impure ferrosilicon. The cooled product consists essentially of a large mass of crystalline alumina. It is crushed, mixed with a ceramic binding material (ball-clay and felspar), moulded, dried, and fired in a porcelain kiln 2

Aluminium peroxide.-When excess of 30 per cent hydrogen peroxide is added to a solution of aluminium hydroxide in 50 per cent potassium hydroxide, a white, amorphous precipitate is obtained, of the composition  $Al_2O_3$ ,  $Al_2O_4$ ,  $10H_2O_5$ , which reacts as a time peroxide. Possibly an initial product  $Al_2O_4$  is formed and partly decomposed by the water

Aluminium hydroxides; aluminates.—Two hydrated oxides of aluminium are found in nature in the crystalline state, namely, diaspore, Al<sub>2</sub>O, H<sub>2</sub>O, which occurs in orthorhombic crystals (holohedial, a b: c = 0.9372 1 1.6038) of density 3.30-3.15, and hydrargillite (or yibbsite), Al<sub>2</sub>O<sub>3</sub> 3H<sub>2</sub>O, which occurs in fibrous, monochine crystals (holohedial, a b c = 1.7089 · 1 · 1.9184,  $\beta$  85° 29') of density 2.42. The most important naturally occurring hydrated oxide of aluminium, however, is bauxite, a white, yellowish, ied, or brown clay-like, amorphous material originally found at Les Beaux near Arles in the south of France. Bauxite varies widely in composition, and consists of amorphous, colloidal, hydrated alumina (with perhaps a little diaspore and hydrargillite) associated with varying amounts of ferric hydroxide, clay, quartz, sand, etc. It is therefore better regarded as a rock than as a mineral 4 Formerly, bauxite was regarded as a mineral of the formula Al<sub>2</sub>O<sub>3</sub>.2H<sub>2</sub>O<sub>5</sub> but most bauxites more nearly approach the ratio Al<sub>2</sub>O<sub>3</sub> H<sub>2</sub>O than Al<sub>2</sub>O<sub>3</sub> 2H<sub>2</sub>O 5 Bauxite is a very valuable source of aluminium, it occurs mainly in the department Var (France), in County Antrim (Ireland), and in the states of Alabama, Georgia, and Arkansas (America).

When excess of ammonium hydroxide is added to an aqueous solution of an aluminium salt, a precipitate is obtained, white, opaque, and amorphous at 100°, transparent and gelatmous at ordinary temperatures. The precipitate has a pronounced tendency to pass into colloidal solution when washed with water. Air-dried in hot weather, its composition's corresponds to the formula Al<sub>2</sub>O<sub>8</sub>.3H<sub>2</sub>O or Al(OH)<sub>3</sub>, dried at 100°, or at the ordinary

Cowles, J. Ind. Eng. Chem., 1913, 5, 331
 O. B. Jacobs, U.S. Pat., 659,926., Higgins, U.S. Pat., 775,654; Williamson and Boeck, J. Ind. Eng. Chem., 1912, 4, 672. See also the following references: Electrochem. Ind., 1903, 1, 15, 1905, 3, 30, 406, Electrochem. and Metall. Ind., 1909, 7, 228; Min. Ind., 1911, 20, 31.
 Terni, Att. R. Accad. Lincei, 1912, [v.], 21, 11, 104

<sup>&</sup>lt;sup>8</sup> Terni, Att. R. Accad. Lincei, 1912, [v.], 21, 11. 104

<sup>4</sup> Bauer, Jahrb. Min., 1898, 11 163; Holland, Geol. Maq, 1903, 59; Dittler and Doolter, Zetsch. Chem. Ind. Kolloide, 1911, 9, 282; Andié Gautiei, Rev. gen. Chim. pure

appl., 1910, 13, 889.

Arsandaux, Compt. rend., 1909, 148, 936, 1115; see also E. Martin, Ann. Chim. anal., 1913, 18, 297.
When precipitated from aluminium sulphate, the hydroxide is contaminated with basic sulphate.

. temperature over concentrated sulphuric acid, the composition is that of a dihydrate, Al<sub>2</sub>O<sub>3</sub> 2H<sub>2</sub>O The amorphous trihydrate is also obtained by heating an alkali aluminate with ammonium chloride, or by boiling basic aluminum carbonate with water, when an alkali aluminate solution is boiled, the tuhydrate slowly separates in a crystalline form 1 A monohydrate, Al<sub>2</sub>O<sub>3</sub>, H<sub>2</sub>O, is said to be obtained by heating amorphous alumina with water m a closed tube to 250°.2

The amorphous mono- and di-hydrates are very hygroscopic substances, absorbing water with the formation of the trihydrate or normal aluminium hydroxide At a red heat all the hydrates are converted into alumina

A colloidal solution of aluminium hydroxide was obtained by Crum from aluminium acctate solution. This was heated to obtain a piccipitate of basic acetate, and the precipitate dissolved in 200 times its weight of boiling water The solution was then maintained at 100° for some days, when complete hydrolysis occurred. The liquid was diluted and heated to 100° until all the acetic acid had been volatilised, fresh water being added from time to time. A colourless, tasteless, neutral solution of aluminium hydroxide was thus obtained, readily coagulated by salts and a number of acids, and the gel thus obtained dissolving only in concentrated acids The solution did not act as a mordant, and when evaporated at 100°, left a residue difficultly soluble in acids 3 Graham obtained aluminium hydroxide in colloidal solution by dialysing a solution of aluminium chloride saturated with The colloidal solution so obtained acted as a mordant, aluminium hydroxide and was readily coagulated by acids, bases, and salts to agel soluble in dilute acids 1 Colloidal aluminium hydroxide exhibits anodic cataphoresis, it is seen to be a suspension when examined in the ultra-microscope

In the absence of salts, freshly precipitated aluminium hydroxide is perceptibly soluble in ainmonia, and much more so in methylamine and other organic bases. 6 It is also readily soluble in acids and alkali hydroxides When kept under water for several months, it becomes difficultly soluble in acids and alkalies, concentrated sulphune acid excepted 7. The naturally occurring hydroxides are not readily attacked by acids

Precipitated aluminium hydroxide assumes a bright red colour, not destroyed by dilute acetic acid, when boiled with water containing a drop or two of I per cent. alizarm solution This test readily distinguishes it from gelatinous hydrated silica. Aluminium hydroxide also forms soluble

E. T. Allen, Chem. News, 1900, 82, 75, Allen and Rogers, Amer. Chem. J., 1900, 24, 804; Cannelley and Walker, Trans. Chem. Soc., 1888, 53, 59, Bonsdorft, Page Annaten, 1838, 27, 275; Cossa, Zeitsch für Chem., 1870, 13, 413, St. Gilles, Ann. Chim. Phys., 1866, [in.], 46-88, Lowe, Zeitsch für Chem., 2, 247, Ditte, Compt. cend., 1893, 116, 183, De Schulten, ibid., 1896, 122, 1127, cf. Ramsay, Trans. Chem. Soc., 1877, in. 395, Russ, Zeitsch. anory. Chem., 1904, 41, 216, Schlumberger, Bull. Soc. chim., 1895, [in.], 13, 41, Maumené, Villiers, ibid., 1895, [in.], 13, 309.
 Mitscherlich, J. prakt Chem., 1861, 83, 468; see also De Senarmont, Compt. rend., 1851, 32, 762.

<sup>&</sup>lt;sup>2</sup> Mitscheilich, J. prakt Chem., 1861, 83, 400; see also Do Golimano, 1861, 32, 762.

<sup>3</sup> Grum, Quart. J. Chem. Soc., 1854, 6, 216, Ann Chim. Phys., 1854, [iii.], 41, 185.

<sup>4</sup> Graham, Phil Tians, 1861, 151, 183, Ann Chim Phys., 1862, [iii.], 65, 175. See also Schlumberger, loc. ci.; W. Biltz, Ber., 1902, 35, 4431, Hantsch and Besch, Annalen, 1902, 323, 30; A. Muller, Zeitsch. anorg. Chem., 1905, 43, 320; 1908, 57, 312, Scheneder, Annalen, 1800, 257, 359, Kawamura, J. Coll. Sci. Tokyo, 1908, 25, vin. 1; Bentley and R. P. Rose, J. Amir. Chem. Soc., 1913, 35, 1490, Rose, Kolloid. Chem. Beihette, 1914, 6, 1.

<sup>5</sup> Hildebraud, J. Amer Chem Soc., 1913, 35, 864.

<sup>6</sup> Renz, Ber., 1903, 36, 27, 51.

<sup>7</sup> Tommasi, Chem. Zenti., 1905, ii. 605; Chem. Soc. Abstr., 1905, 88, ii. 712

complex substances with many organic hydroxy-compounds. Further, it enters into combination with many organic colouring matters, producing coloured, insoluble lakes. I pon this property depends the use of aluminium salts as mordants in dyeing.

The solubility of aluminium hydroxide in acids is due to the fact that it acts as a weak base and reacts with acids to produce aluminum salts. In the same way, the solubility of aluminium hydroxide in alkali hydroxides is attributed to the feeble acidic character of the hydroxide - It is, in fact, an amphoteric hydroxide The minute amount of aluminium hydroxide present in aqueous solution in equilibrium with the solid phase must be supposed to dissociate in two ways -

Al 
$$+3011' \Longrightarrow \Lambda I(OH)_3 \Longrightarrow H' + H_2 \Lambda IO_3'$$
 (or  $\Lambda IO_2'$  and  $II_2O$ ).

The hydroxide is weaker as an acid than as a base, and the affinity constant for the acid dissociation represented above has been estimated to be approximately 1 10 10, i.e. the acid is of the same order of strength as borio acid and its alkali salts must be perceptibly hydrolysed in aqueous solution:

From solutions of alumnium hydroxide in alkali hydroxides a number of solid substances, some of them crystalline, have been isolated, which must be regarded as salts of aluminum hydroxide (aluminic acid). These salts are called aluminates, and are mentioned later (p. 76). The nature of the solutions obtained by dissolving aluminium hydroxide in alkali hydroxides has been the subject of much discussion. The freezing point of a dilute solution of sodnin hydroxide is not changed by dissolving aluminium hydroxide in it. This would be expected if each OII' ion used up leads to the production of one aluminate amon, re the result points very clearly to the presence of a meta-aluminate, NaAlO., in solution. Determinations of the ratio  ${\rm Al}_2{\rm O}_3 \cdot {\rm Na}_2{\rm O}$  (or  ${\rm K}_2{\rm O}$ ) in solutions saturated with aluminium hydroxide are not conclusive, since the above ratio is much greater when aluminium is dissolved m alkah than when alummum hydroxide is dissolved,3 Moreover, unless the atomic ratio Na (or K) to Al exceeds the value two, the solutions are unstable. The equivalent conductivity gradually rises, and aluminium hydroxide is slowly precipitated 1. The change in conductivity is in harmony with the view that the salt of a monobasic acid is undergoing hydrolysis, Moreover, no change in conductivity is observed without the simultaneous deposition of aluminium hydroxide, so that the latter, when produced by hydrolysis, does not first pass into colloidal solution These results negative the suggestion 5 that while some of the aluminium hydroxide dissolves in alkali hydroxides with the formation of alkali aluminates, most of the hydroxide merely passes into solution as a colloidal hydrosol, which, on standing, slowly reverts either to a crystalloidal form or to a colloidal hydrogel, and so

Slade, Zeitsch aning Chem., 1912, 77, 457, Blum, J. Amer. Chem. Soc., 1913, 35, 1499; cf. Wood, Trans Chem. Soc., 1908, 93, 417.
 A. A. Noyes and Whitney, Zeitsch physikal. Chem., 1894, 15, 694
 Herz, Zeitsch. aning. Chem., 1900, 25, 155, Russ, ibid., 1904, 41, 216, Allen and Rogers, Amer. Chem. J., 1909, 24, 301, Heiz, Zeitsch. Elektrochem., 1911, 17, 403; Slade, ibid., 1911, 17, 261, 1912, 18, 1.
 Hantzsch, Zeitsch. aning. Chem., 1902, 30, 206. Russ, loc cd., Slade and Polack, wide wife.

vide unfra 

Mahin, Ingraham, and O. J. Stewart, J. Amer. Chem. Soc., 1913, 35, 30. cf. J. Hildebrand, J. Amer. Chem. Soc., 1913, 35, 864, Blum, abid., 1913, 35, 1499, 1914, 36, 2383;
Mahin, abid., 1914, 36, 2381.

precipitates. A study of the variation of the hydrogen ion concentration in an aluminium chloride solution, as sodium or potassium hydroxide is gradually added to the liquid, has been made by Hildebrand and by Blum.<sup>1</sup>. The general nature of their results may be seen by reference to fig. 12, the points A, B, C are points of inflexion on the curve, and correspond respectively to the commencement of precipitation of the hydroxide, the completion of the precipitation, and the completion of the solution of the precipitate in sodium

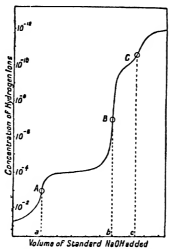


Fig. 12.—Change of hydrogen ion concentration during precipitation of aluminium hydroxide from aluminium chloride and solution of the precipitate in sodium hydroxide.

hydroxide. It is found that bc is one-third of ab, ie the results point clearly to the existence of aluminates NaAlO<sub>2</sub> and KAlO<sub>2</sub> in aqueous solution:—

$$AlCl_3 + 3NaOH = Al(OH)_3 + 3NaCl$$
  
 $Al(OH)_3 + NaOH = NaAlO_2 + 2H_2O$ 

304, are opposed to the view that colloidal solution occurs to any appreciable 1833, in. Moreover, observations with the ultra-microscopic fail to indicate that De Schulttions are suspension-colloids <sup>2</sup> The microscopic evidence, however, Lettsch ano: conclusive.

Zettsch ano; conclusive.

Maumené, Vir wing hydrated aluminates have been obtained in the solid state,

Mitscherhe, a in the crystalline form, by the interaction of aluminium with

Crun, Quart. In hydroxides:—

Graham, Phil 2

<sup>8</sup> Crum, Quart. M Dydroxides:—

Graham, Phil 2,

also Schlumberger, loc. cent., Blum, loc. cet.

1902, 323, 30; A. Muller, ms. Faraday Soc., 1914, 10, 150; Hildebrand, loc. cet.

Annalen, 1890, 257, 359; Kalen and Rogers, Amer. Chem. J., 1900, 24, 304; see also

B. P. Ross, J. Amer Chem. Soc. 907, 29, 300; Fremy, Ann. Chem. Phys., 1844, [11], 12,

Hildebrand, J. Amer Chem. sbid., 1851, [ini.], 33, 13; Mallard, sbid., 1873, [iv.], 28,

Renz, Ber., 1903, 36, 2751.

'2; Beckmann, J. prakt. Chem., 1882, [in.], 26, 385;

Tommasi, Chem. Zenti, 1905, 11 Gaudin, sbid. 1862, 54, 687.

#### ALUMINIUM.

K(AlO<sub>2</sub>).1·5H<sub>2</sub>O Na(AlO<sub>2</sub>).2H<sub>2</sub>O LiH(AlO<sub>2</sub>)<sub>2</sub>.5H<sub>2</sub>O Ba(AlO<sub>2</sub>)<sub>2</sub>.5H<sub>2</sub>O Sr(AlO<sub>2</sub>)<sub>2</sub>.4H<sub>2</sub>O \$

 $\begin{array}{l} {\rm Ca_2Al_2O_5.7H_2O} \\ {\rm Ba_2Al_2O_5.5H_2O} \\ {\rm Tl_4Al_2O_5.7H_2O} \end{array}$ 

Ca<sub>3</sub>(AlO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O Sr<sub>3</sub>(AlO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O Ba<sub>3</sub>(AlO<sub>3</sub>)<sub>2</sub>.7H<sub>2</sub>O

A number of anhydrous, crystalline alluminates occur as minerals; eg:

These four muncials crystallise in regular octahedra, and are isomorphous with magnetite, Fe(FeO<sub>2</sub>)<sub>2</sub>, and chromite, Fe(CrO<sub>2</sub>)<sub>2</sub>. The mineral chrysoberyl,

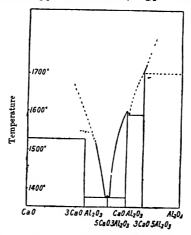


Fig. 13.—Equilibrium diagram for the system lime - alumina.

or beryllium aluminate,  $Bc(AlO_2)_2$ , is not isomorphous with the preceding, but crystallises in the orthorhombic system (a b c=0 470; 1.0.580) and is isomorphous with olivine,  $Mg_2SiO_4$  Magnesia spinel is often found in beautiful red crystals, which are used as genistones, such crystals contain a little chromic oxide Beautiful yellowish-green crystals of chrysoberyl, found in Ceylon, are also used as genistones. The aluminates of barium, calcium, beryllium, magnesium, zinc, manganese, and cobalt were prepared in the crystalline form by Ebelmen by intensely heating alumina and the requisite oxide with boron sesqui-oxide until the latter substance, which initially acts as a solvent, had been largely volatilised <sup>1</sup> The blue mass obtained in qualitative analysis when testing on charcoal for aluminium contains cobalt aluminate.

<sup>&</sup>lt;sup>1</sup> Ebelmen, Ann Chim. Phys. 1848, [111], 22, 211, 1851, [111], 33, 34, Deville and Caron, ibid., 1865 [1v], 5, 104, Daubric, Compt. rend., 1854, 39, 135; Dufau, J. Pharm. Chim., 1901, [v1.], 14, 25; Hedvall, Arkiv. Kem. Min. Geol., 1914, 5, No. 6 (Co), Jaeger and Simek, Proc. K. Akad. Wetensch. Amsterdam, 1914, 17, 239, 251 (Li).

A thermal study of the system lime - alumina has shown that four anhydrous calcium aluminates can be obtained, of the formulæ  $3\text{CaO.Al}_2\text{O}_3$ ,  $6\text{CaO.3Al}_2\text{O}_3$ , and  $3\text{CaO.5Al}_2\text{O}_3$ . The second and third melt at  $1587^\circ$  and  $1387^\circ$  respectively, the others have no melting-point. The third and fourth compounds are dimorphous  $^{1,2}$ . The nature of the equilibrium diagram is indicated in fig. 13. Only one magnesium aluminate has been obtained, namely, spinel,  $MgO.Al_2O_3$ ,  $^{1,2}$ .

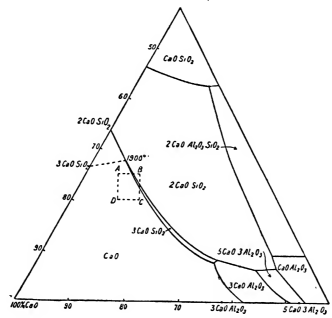


Fig. 14.—Projection of liquidus surface for the system lime - silica - alumina.

Tricalcium aluminate, 3CaO.Al<sub>2</sub>O<sub>2</sub>, is an important constituent of Portland cement clinker, which consists mainly (over 90 per cent ) of silica, lime, and alumina. Its constitution has long been the subject of discussion. A complete thermal and optical study of the system lime - silica - alumina has been made by Rankin and Wright,3 and the results, so far as they bear on the constitution of Portland cement, are shown in fig 14. Considering only the lime, silica, and alumina content, the variations in the composition of the commercial cement are restricted to the area ABCD. It may be deduced from the diagram by aid of the Phase Rule that a fluid mass of this composition, when slowly cooled until complete solidification has occurred, is converted

Shepherd, Rankin, and Wright, Amer. J. Sci., 1945, [iv], 28, 293.
 Rankin and Merwin, J. Amer. Chem. Soc., 1916, 38, 568.
 Rankin and Wright, Amer. J. Sci., 1915, [iv.], 39, 1; J. Shepherd, Rankin, and Wright, J. Ind. Eng. Chem., 1911, 3, 211; Rankin, ibid., 1915, 7, 466.

into a mixture of tricalcium silicate, 3CaO.SiO, dicalcium silicate, 2CaO.SiO, and tricalcium aluminate, 3CaO.Al2O3.

Portland cement clinker should therefore consist essentially of these three compounds if, in its manufacture, the heating be prolonged for a length of time sufficient for equilibrium to be established. This deduction has been verified by experiments carried out in the laboratories of the United States Bureau of Standards Since, however, complete equilibrium is not reached in its manufacture, the cement also contains hime, CaO, and the aluminate  $5{\rm CaO}~3{\rm Al}_2{\rm O}_3$  as minor constituents. The non-oxide contained in the cement may be present as calcium ferrite or magnetice, or hold in solid solution in the compounds 2CaO SiO, and 5CaO 2Al<sub>2</sub>O<sub>3</sub>, the magnesia and alkalies are apparently held in solid solution in the compounds 2CaO SiO2 and 3CaO.Al<sub>2</sub>O<sub>8</sub>1

In contact with water, the compounds CaO Al<sub>2</sub>O<sub>3</sub>, 3CaO Al<sub>2</sub>O<sub>3</sub>, and 5CaO 3Al2O3 yield an amorphous hydrate, 3CaO Al2O3 all2O, which in time crystallises, except with tricalcium aluminate, amorphous aluminium hydroxide is also formed. When hydrated with aqueous calcium sulphate an additional compound, calcium "sulpho-aluminate," 3CaO Al<sub>2</sub>O<sub>3</sub> 3CaSO<sub>4</sub>,cH<sub>2</sub>O<sub>5</sub>

is formed, which crystallises in long prismatic needles 2

Sodium aluminate is manufactured from bauxite, which is roughly powdered, calcined, and introduced into sodium hydroxide solution of specific gravity 1°15 The mixture is heated under high-pressure steam (80 lbs. per sq. mch), when the alumna passes into solution. The liquid is passed through a filter-press, and then through wood-pulp to free it from ferric hydroxide, etc. Aluminium hydroxide is prepared from this solution by stirring into it a quantity of the hydroxide previously prepared. On standing, the greater part of the aluminum separates out from the solution as aluminum hydroxide. The precipitate is washed in a filter press, and the mother liquor concentrated and used again for attacking more bauxite. This process, due to Bayer, has largely superseded the older method of decomposing the sodium aluminate by passing carbon dioxide through the solution

Aluminium sesqui-sulphide, Al.S., was first obtained by Fremy by heating alumina to redness in a current of carbon disalphide vapour. It may be obtained by the action of hydrogen sulplude upon alumina at a red heat or by the action of sulphin on a heated maxture of alumma and carbon 3

Aluminium sulphide is readily prepared by heating alumina with galena in a muffle furnace, or by the direct union of its elements. The powdered elements are mixed in a friedly crucible, heated till the sulphur fuses, and cooled The reaction is then brought about as in a thermit reaction by means of a firing powder ignited by a piece of magnesium ribbon  $^{4}$  . The sulplinde is

Chem., 1911, 71, 182.

<sup>1</sup> P. H. Bates, Concrete-Cowent Age (Conent Mill Section), 1913, 2, 3, 1914, 4, 29, 38; 1915, 7, 1, A. A. Klein and A. J. Plulips, Lighth Into Cong. Appl. Chem., 1912, 5, 73; cf. Janecke, Zeitsch. anny Chem., 1911, 73, 200, 1912, 74, 428, 76, 357, 1914, 89, 355; 1916, 93, 271. and Kuhl, Tonind. Zev., 1914, 38, 365, who consider that a compound Scao. Al<sub>2</sub>O<sub>3</sub> 28iO<sub>2</sub> exists in the cement.

A. A. Klein and A. J. Plulips, Tech. Papers Bur. Stand., 1914, No. 43, Trans. Amer.

Cer. Soc., 1914, 16, 313; J. Wesh. Acad. Sec., 1914, 4, 57;

3 Gautter, Compt. rend., 1906, 143, 7, Bucheri, Zeitsch. angeu. Chem., 1892, 5, 483.

Early references. Frémy, Ann. Chem. Phys., 1853, [m.], 38, 322. Salastici, ibid., 1881, [v.], 22, 88, Vincent, Phil. Mag., 1857, [n.], 14, 127. Spring, Bir., 1883, 16, 1011; Rechel, J., makt. Chem., 1875, 117, 12, 55, Stein, ibid., 1871, [n.], 3, 13, Knopp and Ebell, Dingl. Poly. J., 1878, 229, 69, 173.

4 Fonzes-Diacon, Compt. rend., 1900, 130, 1314, W. Biltz and Caspan, Zeitsch. anny. Chem., 1911, 71, 182.

purified by sublimation in vacuo at 1100° to 1250°, or in mitrogen at 1500° , to 1600°.

According to Biltz and Caspari, aluminium sulphide crystallises in white, asbestos-like needles (which are possibly hexagonal) and forms mixed crystals with alumina. It melts at 1100° ± 10° and its density at 13° is 2.02. Heated to 2100° in an inert atmosphere it loses sulphur and forms aluminium subsulphide, AlS., heated in air or oxygen it is converted into alumina and sulphur dioxide.

Aluminium sesqui-sulphide is rapidly and completely decomposed by water with the formation of aluminium hydroxide and hydrogen sulphide, a behaviour in accordance with the fact that it is the salt of a very weak, practically insoluble base and a very weak acid. The decomposition of the monosulphide by hydrochloric acid is represented by the equation:-

$$2AIS + 6HCl = 2AICl_3 + 2H_2S + H_2$$

By heating a mixture of alumina and a carbonate to redness in a current of carbon disulphide vapour, the following double sulphides have been obtained: 2 potassium-aluminium sulphide, 3K<sub>2</sub>S, Al<sub>2</sub>S<sub>3</sub>; sodium-aluminium sulphide, 3Na<sub>2</sub>S, Al<sub>2</sub>S<sub>3</sub>, and barium-, strontium, and calcium-aluminium sulphides. Aluminium silver sulphide, 4Ag, 55Al, S, melts at 1035°.3

Aluminium selenide, Al<sub>2</sub>So<sub>3</sub>, may be prepared from its elements as described for the sulphide, or by reducing lead sulphiate by means of aluminium.<sup>4</sup> It is a greyish-yellow solid of density 3 437, and is readily decomposed by water with the evolution of hydrogen sclenide.

Aluminium telluride, Al, Te, 1s prepared by the direct union of its elements.<sup>5</sup> It is a grey solid, decomposed by water with the evolution of hydrogen telluride, which, however, readily decomposes

Aluminium sulphite.—Various basic salts have been described.6 The acid sulphite solution has been used for purifying beet sugar 7

Aluminium sulphate, Al<sub>2</sub>(SO<sub>1</sub>), is prepared in the anhydrous state by heating the crystalline, hydrated salt. The latter melts in its water of crystallisation, swells up, and eventually leaves a porous, white residue of anhydrous sulphate.

The authydious sulphate has a density of 2713 at 17°, and its specific heat (0° to 100°) is 0 1855.8 At a red heat it decomposes, leaving a residue of alumina; decomposition becomes appreciable at 770°.0 It dissolves slowly

A solution of aluminium sulphate is readily prepared by dissolving aluminium hydroxide in dilute sulphuric acid. The solution crystallises with difficulty, the hydrate Al2(SO1)3.18H2O being deposited in thin, six-

*(*%,

<sup>1</sup> Cf. Mourlot, Compt. rend., 1896, 123, 54.

3 Jaeningen, Chem. Zentr., 1895, 11. 205

5 Cambi, Att. R. Accad. Linces, 1912, [v.], 21, 11. 837; for other double sulphides, see Vols. VII., VIII., IX., or Houdard, Compt. rend., 1907, 144, 801, 1114.

5 Fonzes-Diacon, loc. cit.

6 Fonzes-Diacon, loc. cit., Whitehead, J. Amer. Chem. Soc., 1895, 17, 849.

6 Gougginsperg, Annalen, 1843, 45, 132; Muspratt, ibid., 1844, 50, 259, Scutari and Manzoni, Gazzetta, 1884, 14, 360, Scubert and Eltern, Zertsch. anny. Chem., 1893, 4, 44.

7 Becker, Dingl. poly. J., 1835, 257, 300.

8 Nilson and Pettersson, Ber., 1880, 13, 1459, Compt. rend., 1880, 91, 232; cy. Favre and Valson, ibid., 1872, 75, 803.

9 Friederich, Centr. Min., 1912, pp. 174, 207.

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sided, nacreous plates. This hydrate has also been obtained in the form of tetrahedra. At low temperatures the hydrate Al<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub> 27H<sub>2</sub>O separates in trigonal crystals (a c=1 0.5408). Other hydrates, with 16H<sub>2</sub>O, 12H<sub>2</sub>O, 10H<sub>2</sub>O, 9H<sub>2</sub>O, 6H<sub>2</sub>O, 3H<sub>2</sub>O, and 2H<sub>2</sub>O have been described. The hydrates with 9H<sub>2</sub>O and 10H<sub>2</sub>O are said to be precipitated by alcohol, and to absorb water from a damp atmosphere, forming the hydrate with 18H2O hexahydrate results from the action of concentrated sulphuric acid on the hydrate with 18H2O,2 the trihydrate forms regular tetrahedra 3. The system

aluminium sulphate - water has not yet been systematically investigated 4. The hydrate Al<sub>2</sub>(SO<sub>1</sub>), 18H<sub>2</sub>O is practically investigated in alcohol. Its density is 1 69, and its specific heat (15° to 52°) is 0 353 As a white, fibrous offlorescence on shale and other rocks, this hydrate occurs as the mineral

alunouen.

;,•:•

Aluminium sulphate is used in paper making, as a mordant, and for the cipitation of sewage. The crude, very impure commercial preparations precipitation of sewage are manufactured by heating powdered, calemed china-clay or banxite with sulphuric acid, the product being caused to solidify into blocks aluminium sulphate is required free from iron salts it is prepared from aluminum hydroxide, made as described previously (p. 79), and sulphuric acid It is impossible to separate ferrie sulphate from aluminium sulphate by crystallisation, although the two sulphates form neither a compound nor mixed crystals, but commercial aluminium sulphate may be freed from iron by crystallisation if the non sulphate present is reduced to the ferious state by hydrogen sulphide, sulphuric acid, etc 5
The acid sulphate Al<sub>2</sub>(SO<sub>4</sub>), IL<sub>2</sub>SO<sub>4</sub> 3H<sub>2</sub>O has been prepared by Band, and

also by With, while Silberberger has described an acid sulphate of the composition Al<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>.3H<sub>2</sub>SO<sub>4</sub>.7H<sub>2</sub>O<sup>4</sup> When aluminium sulphate is boiled with concentrated hydrochloric acid and the liquid cooled, crystals of aluminium chlorosulphate, AlCISO, 6H,O are deposited, which are decomposed by water.

About twenty different basic sulphates of aluminium have been described, but beyond the statement that the substance Al<sub>2</sub>O<sub>1</sub> 2SO<sub>1</sub> and one or two hydrates of it are probably definite compounds, practically nothing can be said about them. A hydrated basic salt of the composition Al<sub>2</sub>O<sub>3</sub> SO<sub>3</sub>9H<sub>2</sub>O occurs naturally as the inmeral websterite

Alums .- Aluminium sulphate forms double salts with the sulphates of sodium, potassium, rubidium, casium, ammonium, and thallium (thallous sulphate), of the type R'SO<sub>4</sub>Al<sub>2</sub>(SO<sub>4</sub>), 24H<sub>2</sub>O. These double salts are called alums, the salt Cs<sub>2</sub>SO<sub>1</sub>Al<sub>2</sub>(SO<sub>1</sub>)<sub>3</sub> 24H<sub>2</sub>O, for example, being called cusium

<sup>&</sup>lt;sup>1</sup> Gawalowski, Chem Zentr., 1906, ii. 1230.

Schmatolla, Zeitsch. angro. Chem., 1903, 16, 202.
 Boutzoureano, Ann. Chim. Phys., 1889, [vi.], 18, 289.
 See Kremann and Huttinger, Jahrb. K. K. Rewhsanstalt, 1908, 58, 637.
 Wirth and Bakke, Zeitsch. anorg. Chem., 1914, 87, 47, cf. Witth, Zeitsch. angew.

Wirth and Bakke, Zeitsch. anoig. Chem., 1913, 87, 41, cf. Willin, Zeitsch. anoig. Chem., 1913, 26, 81.
 Baud, Compt. end., 1903, 137, 492, Wirth, Zeitsch. anoig. Chem., 1913, 79, 380;
 Silberberger, Monatsh., 1904, 25, 220.
 Recoura, Compt. rend., 1902, 135, 736
 For Al<sub>2</sub>O<sub>2</sub>2SO<sub>2</sub> and hydrates, see Spence, D.R.P., 167,419 (1903); Kremann and Huttinger, loc. etc. Other references are. Mans, Poyg. Annalen, 1827, 11, 80; Rammelserg, ibid., 1383, 42, 583, Bley, Annalen, 1854, 89, 174; Bottinger, lod., 1888, 244, 224; Lowe, J. prakt. Chem., 1860, 79, 423, Marguente, Compt. rend., 1880, 90, 1354; Athanassesco, ibid., 1886, 103, 271; Debray, Bull. Soc. chim., 1867, [n.], 7, 9, Gudumer, Chem. Zentr., 1895, n. 1087, Pickering, Trans. Chem. Soc., 1907, 91, 1981.

aluminium alum, or simply casium alum. All these salts are isomorphous with one another, usually crystallising in regular octahedra or combinations of the octahedron and cube. Further, isomorphous series of double sulphates of the same type,  $\rm\,R_2^iSO_1, R_2^m(SO_1),\,24H_2O,\,are$  known, in which  $\rm\,R^i$  may, as before, be Na, K, Rb, Cs, NII, or Tl,2 but m which Rin is not Al, but Ga, In, Cr, Fc, V, Co, Ti, Mn, or Rh. All these salts I crystallise in the regular system in crystals of octahedral habit, and, like the double aluminium sulphates, are also called alums In naming one of these alums, both metals must be mentioned, the salt Rb<sub>2</sub>SO<sub>1</sub>,Co<sub>2</sub>(SO<sub>1</sub>),  $24H_2O$ , for example, being called *rubidium* cobalt alum, while with an aluminium alum it is usual to mention only the univalent metal present. Finally, analogous and isomorphous double selevates are known, and called selenium alums, eg, the salt Tl, ScO, Cr, (ScO,), 21H,O is called thallium chromium selenium alum, and mixed alums, in which one component salt is a sulphate and the other a scienate, have also been prepared.1

The alums are described in this series of text-books under the heading of the tervalent metals present. Hence only the aluminum alums are described in this chapter.

The alums are readily prepared in the crystalline form by allowing aqueous solutions of the requisite sulphates, mixed in the ratio of their molecular weights, to evaporate slowly at the ordinary temperature crystals which then separate almost invariably possess an octahedral habit, but in the presence of small quantities of foreign substances, the habit may undergo considerable modification? Thus, it has long been known that below 45°, and in the presence of a trace of basic alum, potassium alum crystallises in cubes and not in octahedra 6. Moreover, it is an old observation that sodium alum may separate out in monoclinic crystals,7 but according to Surgunoff the monoclinic form (a  $b = 2510 - 1 \cdot 0.908$ ,  $\beta = 109^{\circ}$  1') has the formula Na<sub>2</sub>SO<sub>4</sub>Al<sub>2</sub>(SO<sub>4</sub>), 22H<sub>2</sub>O and separates from supersaturated solutions at temperatures above 20.8

The densities of the aluminium alums at 18° to 20°, according to the careful determinations of Pettersson, are as follows -9

| Alum .    | Na    | K     | Rb    | ('s   | XII, | Tl    |
|-----------|-------|-------|-------|-------|------|-------|
| Density . | 1:691 | 1.751 | 1.890 | 1 997 | 1662 | 2 370 |

<sup>&</sup>lt;sup>1</sup> Lithium alum was said by Kratovanszky to be obtained from aqueous solution when crystallisation takes place below 11°C (*Schurugger's J.*, 1828, 54, 349). Its existence was defined by Rammelsberg (*Pharm Centr.*, 1849, p. 106). Schreinomaker and de Waal (*Chem. Weekblad.*, 1906, 3, 539) have shown that hthium alum does not exist at 30° A study of the system Li  $SO_4 - Al_2(SO_4)_3 - H_2O$  at 0°C, would probably prove of interest <sup>2</sup> The sulphates of hydroxylamine and numerous alphatic amines also combine with the sulphates of alumnum star, to turn along

sulphates of aluminium, etc., to form alums.

All the possible combinations are not known. In a small number of cases the attempted preparation has failed, and in other cases the combinations have not been tried

Gerichten, Annalen, 1873, 168, 214.
 Weber, Pogg. Annalen, 1860, 109, 379, Januetaz, Bull. Soc. chim, 1870, [n], 13, 3;
 Leooq de Boisbaudran, Bull. Soc. chim., 1879, [n.], 31, 290, Weyberg, Chem. Zentr., 1910, n. 1026.

<sup>1910,</sup> n. 1026.
Frémy and Pelouze, Leblanc, Kopp, Annalen, 1855, 94, 122, Von Hauer, Jahresber.,
1865, p. 181; Loewel, Ann. Chim. Phys., 1855, [ni.], 43, 114, Persoz, Ann. Chim. Phys.,
1849, [ni.], 25, 257, Polis, Ber., 1880, 13, 360.
Soret, Arch. Sci. phys. nat., 1881, [ni.], 11, 62
Surgunoff, Bull. Acad. Sci. Petrograd, 1909, p. 1057, Soret suggested 23H<sub>2</sub>O
Pettonson, Nova Acta Soc. Upsala, 1873, [ni.], 9, No. 4, cf. Spring, Bull. Acad. roy.
Belg., 1883, [ni.], 6, 507.

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while the refractive indices for the D-line are given by Soret 1 as follows:-

Na 1 4388 NH, 1.4586 1:4975 1.4564 1.4566 14594

The solubilities of the alums  $^{2}$  are given in the accompanying table  $-^{8}$ 

| Tempera-<br>ture °C.                                   | ن Sodium.                               |                                                                                                     | Potassium                                                                                                      |                                                                                        | Rubidinin,           |                                                                                                       | ('æsııım |                                                              | Ammonium                                                               |                                                                        | Thallium.                                                                  |  |
|--------------------------------------------------------|-----------------------------------------|-----------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|----------------------|-------------------------------------------------------------------------------------------------------|----------|--------------------------------------------------------------|------------------------------------------------------------------------|------------------------------------------------------------------------|----------------------------------------------------------------------------|--|
| Tem v                                                  | В                                       | Λ                                                                                                   | В.                                                                                                             | А.                                                                                     | В                    | Λ.                                                                                                    | В.       | Λ.                                                           | В                                                                      | Α.                                                                     | В,                                                                         |  |
| 0 5 36 15 38 20 40 25 11 30 15 40 50 80 90 92 5 95 100 | 7 111 3 , 9 121 4 ; 3   131 8 8   146 3 | 3.0<br>3.5<br>4.0<br>5.0<br>5.9<br>7.23<br>8.39<br>11.70<br>21.75<br>40.0<br>71.0<br>109.0<br>119.0 | 5 65<br>6 62<br>7 60<br>9 59<br>11 40<br>14 14<br>16 58<br>23 83<br>36 40<br>57 35<br>110 5<br>321 3<br>2275 0 | 0 72<br>0 86<br>1 05<br>1 28<br>1 50<br>1 80<br>2 20<br>3 25<br>7 10<br>12 40<br>21 60 | 3 12<br>3 82<br>5 69 | 0 21<br>0·25<br>0 30<br>0 35<br>0 40<br>0·50<br>0 60<br>0 85<br>1 30<br>2 00<br>3 20<br>5 40<br>10 50 | 18 11    | 21<br>35<br>50<br>63<br>77<br>92<br>109<br>119<br>201<br>267 | 3.9<br>9.5<br>12.7<br>15.1<br>19.2<br>22.0<br>30.9<br>41.1<br>66.7<br> | 3·15<br>3 80<br>4 60<br>6·40<br>7 60<br>9 38<br>11 40<br>22 0<br>35 36 | 4·84<br>5·86<br>7·12<br>10·00<br>11·95<br>14·89<br>23·57<br>38·41<br>66·19 |  |

It will be noticed that rubidium and caesium alums are only sparingly soluble in water. The alums are insoluble in alcehol. They melt in their water of crystallisation at the following temperatures (Locke) .-

The alums are completely dehydrated at 200°, information concerning lower hydrates is not very definite. The porous mass obtained by dehydrating potassium alum is called burnt alum, and dissolves slowly but completely in water The ignition of pure ammonium alum leaves a residue of pure alumina, which is conveniently prepared in this manner

Aqueous solutions of the alums have an acid reaction and in dilute

Soret, Arch Sci. phys. nat., 1881, [iii], 12, 553., see also ibid., 1885, [iii], 13, 5; 14, 96., 1883, 20, 520.
 A = grains anhydrous salt per 100 grains water. B = grains crystalline, hydrated salt per 100 grains (extra) water. The solubility of ammonium alum is in need of redetermination. Poggiale, Ann Chim. Phys., 1843, [iii], 8, 467.
 Setterberg, Annalem, 1882, 211, 104; Locke, Amer. Chim. J., 1901, 26, 166.
 Marino, Gizzalla, 1905, 35, ii. 351. Earl of Berkeley, Phil. Trans., 1904, A, 203–189.
 Wadmote, Proc. Chem. Soc., 1905, 21, 150; Hart and Huselton, J. Amer. Chem. Soc., 1914, 36, 2082.
 Sedell, Solubilities (Crosby, Lockwood & Son, 1907).
 Precht and Kraut, Annalem, 1875, 177, 144.
 Losco ut and Mathurm, Bull. Soc. divin., 1888, [ii.], 50, 33; Maumené, ibid., 1886, [ii.], 46, 261, 807.
 Borssien, ibid., 1887, [ii.], 47, 494.
 Naumann, Ber., 1875, 8, 1630., 1877, 10, 457, Muller-Erbach, Ber., 1888, 21, 2222, 1889, 22, 3151; Impton. Chem. News., 1871, 30, 272.
 Favie and Valson, Compt. rend., 1872, 75, 803.
 Locke, vide supra.
 Earl of Berkeley, vide supra.

solution behave as would be expected of solutions of mixed sulphates which did not interact to any appreciable extent to produce complexes.

Sodium alum, Na<sub>2</sub>SO<sub>1</sub>.Al<sub>2</sub>(SO<sub>1</sub>)<sub>3</sub>.24H<sub>2</sub>O —The existence of this alum, occasionally denied, has been proved by Wadinore and by W. R. Smith <sup>2</sup> The alum is extremely soluble in water. Several methods of preparation have been patented. Sodium alum occurs as the mineral mendonte in South America and Japan.4

Potassium alum, K2SO1 Al2(SO4)3 24H2O.—This alum is an article of commerce, being used as a mordant, in the production of other aluminium mordants, in the manufacture of lake pigments, in dressing skins to produce white leather, in sizing paper, and in the production of fire-proofing materials. To a small extent it occurs naturally, and it is manufactured from aluminium sulphate (prepared from banvite or china-clay) and potassium sulphate, from alum-rock and from alum-shale.

Natural potassium alum occurs as fibrous crystals or as an efflorescence on aluminous minerals at Whitby, Campsie, etc. At Solfatara, near Naples, and in the mlands of Volcano and Milo it occurs in some quantity, and when twice recrystallised from water furnishes very good alum.

The manufacture of alum from alum-rock or alunite is an industry which dates from very early times - It was introduced into Europe in the thirteenth century, and several alum works were established during the fifteenth century, notably the celebrated works at La Tolfa near Civita Vecchia. In this district the manufacture of the so-called Roman alum is still an important industry, and the alum prepared there has always been highly valued on account of its purity. Alunite may be regarded as a double salt of potassium sulphate and a basic aluminium sulphate, K2SO1.Al2(SO1)3 4Al(Oll)3 It is a mineral which occurs in large quantities at La Tolfa, in Hungary, at Puyde-Sancy and Madria in Auvergue, and in numerous other localities, having been formed by the action of volcame gases upon felspathic trachyte. In the modern process, alumite is calcined at 500° to 1000° and the product treated with sulphuric acid A solution is thus obtained which deposits alum when crystallised, leaving an excess of aluminium sulphate in the mother liquors. It is usual, therefore, before crystallisation, to add sufficient potassium sulphate to enable all the aluminium sulphate present to be converted into alum.

The production of alum from alum-shale, an industry of great antiquity, is at the present time only of slight commercial importance. a kind of shale or slate containing iron pyrites disseminated throughout its mass in a very finely divided state. On prolonged exposure to the weather the pyrites undergoes oxidation and the sulphuric acid produced attacks the clay. The weathered product, when leached with water, gives a solution contaming aluminium sulphate and other substances It was known to Agricola and Libavius that this solution would not crystallise well unless an alkali had been added to it, and each of these writers describes the early practice of

<sup>31, 245.</sup> Eng. Pat., 5650 (1881); 5644 (1900), D.R.P., 50,323; 141,670.

adding decomposed urine to the solution to facilitate its crystallisation. Hence the alum prepared must have been mainly ammonium alum. Later, the urine was replaced by potash in the manufacture of alum. The fact that alum contains two bases, alumina and potash, was discovered by Marggrat and emphasised by Lavoisier, but was not generally accepted until 1797, when Chaptal and Vanquelm showed that potash was an essential constituent of ordinary alum, but could be replaced by ammonia, and that alum could be prepared from alumte without the addition of potash because potash was already present in the nimeral. It should be mentioned that the presence of potash in alum was known to Bergman and Scheele, but they looked upon

it only as an impurity 1

The alum-shales at Whithy have been largely worked for alum since the time of Queen Elizabeth, though not worked at the present time. The shales found in the West Riding of Yorkshire, however, are still utilised to a small extent. Unless sufficiently bituminous, the shales are mixed with fuel and slowly roasted, but some shales contain sufficient carbonaceous matter to render the addition of fuel unnecessary. The roasting lasts for ten days, and the temperature never reaches a red heat. The product, which is light red, soft, and porous, is treated with sulphinic acid (density 1.35) at 110° in lead-lined vessels. The solution of impure aluminium sulphate thus obtained is mixed with potassium sulphate and rapidly cooled with sturing to produce small crystals of "alum meal". The meal is dramed, washed with mother liquor from "block alum," dissolved to form a hot, saturated solution, and treated in leaden vessels with a little size, which precipitates a quantity of insoluble matter. The clear solution is then run into tubs fitted with movable, lead-lined staves. After several days the staves are removed and a hole is bored in each mass of "block alum" to allow the mother liquor to drain away. The "block alum" when broken up is ready for the market. When shale is employed containing much non, it is the custom to add potassium chloride, either alone or mixed with sulphate, instead of adding potassium sulphate to the solution of aluminium sulphate. By this means contamination of the alum with isomorphous non alum is prevented, the iron remaining behind in solution as ferric chloride 2

Potassium alum becomes white and opaque on the surface when exposed to the air, a change brought about by the absorption of ammonia and formation of a basic salt. When dried over sulphune acid or heated to 61°, alum loses

18 molecules of water of crystallisation.

When an alkali is slowly added to a solution of alum, a piccipitate is produced which redissolves on sturing until a certain amount of alkali has been added, after which the further addition of alkali leads to the production of a permanent precipitate The solution which is on the point of yielding a permanent precipitate is known in commerce as neutral alum, and is used in dyeing, since it readily gives up alumma to the colouring matter. When heated to 40°, a precipitate is produced the composition of which may be represented by the formula  $K_2SO_4.\Lambda l_c(SO_4)_3$   $4\Lambda l(OH)_3$ , i.e. it is identical in composition with alunite. The precipitate may be obtained in the crystalline form by heating the solution in a scaled tube to 230°.

vol. ii. p. 720.

Further details of the manufacture of alum will be found in Thorpe, A Dictionary of Applied Chemistry (Longmans & Co., 2nd ed., 1912-13), vol. 1., article "Aluminium."

<sup>1</sup> Roscoe and Schorlemmer, A Treatise on Chemistry (Macmillan & Co., 3rd ed., 1907),

Crystals of alum having a cubic habit are obtained by the crystallisation of an aqueous solution of the salt to which a little alkah has been added (p 82) A study of the crystallisation of alum from hydrochloric acid solutions of different concentrations, and at various temperatures, has shown that alum crystals occasionally exhibit the faces of the pentagonal dodecahedron (210) 1

Potassum aluminum sulphate octahydrate, K2SO4Al2(SO4), 811,0, separates in slender crystals when alum is inclted in its water of crystallisation and the fused mass maintained at about 86°. Analogous salts are known containing indium, thallium, and the rare earth metals in the place of aluminium.2

Rubidium alum,  $^3$  Rb,  $SO_1$   $M_2(SO_1)_3$   $24H_2O_5$  and cæsium alum,  $^4$ Cs<sub>2</sub>SO<sub>4</sub> Al<sub>2</sub>(SO<sub>4</sub>), 2411,O, have been prepared by several chemists

Ammonium alum, (NII<sub>1</sub>)<sub>2</sub>SO<sub>1</sub>.Al<sub>2</sub>(SO<sub>1</sub>)<sub>3</sub> 24H<sub>2</sub>O, was formerly manufactured in England to a very large extent, but at the present time it is mainly the potassium salt that is prepared. Commercial alum is often a mixture of potassium and ammonium alums 5

Ammonium aluminium sulphate octahydrate, (NH1),SO1 AL(SO4), 8H2O, may be prepared like the corresponding potassium salt (Marino), or by seeding a supersaturated solution of ammonium alum with a crystal of the corresponding thallie salt,  $(NH_1)_2SO_4$   $Tl_2(SO_1)_3$   $8H_2O_5$ 

Hydroxylamine alum,  $(\tilde{N}\Pi_1^{\dagger}\tilde{O}\Pi)_2\tilde{S}O_4^*\tilde{A}I_2(\tilde{S}O_1)_3$   $24\Pi_2O_6$  has also been

Silver alum, Ag<sub>2</sub>SO<sub>1</sub>Al<sub>2</sub>(SO<sub>1</sub>), 24H<sub>2</sub>O, is said to be prepared by heating silver sulphate and aliminium sulphate, together with a little water, in a sealed tube, and allowing the solution to cool 8. The existence of this alum is denied by Retgers

Pseudo-alums -- A number of double sulphates of the type R"SO<sub>1</sub>-Al<sub>2</sub>(SO<sub>1</sub>), 2411,O have been described, namely the Fe, Zn, Mg, and Mn salts. They are called pseudo-alums, and are not isomorphous with ordinary Very little is known concerning these salts, which require to be reinvestigated.9

Aluminium dithionate,  $Al_2(S_2O_6)_1 18H_2O$ , forms deliquescent plates, soluble in water and in alcohol. A double salt,  $Al_2(S_2O_6)_1 (NH_1)_2 S_2O_6 (27H_2O_6)$ is also known.10

Zemjatschensky, Zutsch. Kryst. Min., 1913, 52, 601, see also Schubnikow, Bull. Acad. Ser. Petrograd, 1913, p. 817, Weber, Peng. Annalen. 1860, 109-379. Lecon. de Borsbaudran, Compt. rend., 1879, 88, 360; L. Wulff, Zeitsch. Kryst. Min., 1881, 5, 81, N. S. Kurnakoff, thad., 1881, 5, 591
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<sup>The references exted in the preceding toothote, and Hattanu Huserton, J. Amer. Soc., 1914, 36, 2082.
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Kluss, Annalen, 1888, 246, 179, 284.</sup> 

87

Aluminium selenite, Al<sub>2</sub>(SeO<sub>1</sub>)<sub>3</sub> 7H<sub>2</sub>O, is produced by treating the basic selenite obtained from aluminium sulphate and sodium selenite with a dilute solution of selemous acid. It is sparingly soluble in water. Two acid selemites of the formula 241,(SeO<sub>3</sub>), 3H<sub>2</sub>SeO<sub>3</sub>, 9H<sub>2</sub>O and Al<sub>2</sub>(SeO<sub>3</sub>)<sub>1</sub> 3H<sub>2</sub>SeO<sub>4</sub> 2H<sub>2</sub>O have also been described <sup>1</sup>

Aluminium selenate, Al (SeO1), is formed by dissolving aluminium hydroxide in selenic acid and heating to expel the excess of acid. It is

soluble in water, but no hydrates have been described

Aluminium selenate forms double selenates with the selenates of sodium, potassium, rubidium, caesium, ammonium, and thallium the formulæ of which are of the type R<sub>3</sub>SeO<sub>4</sub> Al<sub>2</sub>(SeO<sub>4</sub>)<sub>3</sub> 24H<sub>2</sub>O, i.e. they are selenium alums (p. 82). They have been examined by various chemists? The densities of the selenium alums are as follows (Pettersson) -

Aluminium chromate. 1 yellow, flocculent precipitate of basic aluminium chromate, Al<sub>2</sub>O<sub>p</sub>CrO<sub>1</sub>7H<sub>2</sub>O<sub>r</sub> is produced by adding potassium chromate to a solution of alum.

Aluminium molybdate. A white, amorphous precipitate of the composition 5Al\_O\_2MoO\_3/H\_O is obtained by adding sodium molybdate, Na\_MoO\_5 to a solution of alum. The double salts Al\_2(Mo\_2O\_7)\_r 3 \cdot 2 Mo\_2O\_7. where X is Na, K, or NII, are known and possess 22, 20, and 20 molecules of water of crystallisation respectively.4

Aluminium silicomolybdate, 2M2O3 (SiO2 12MoO3)3 93H2O, crystallises in yellow, regular octahedra, isomorphous with the corresponding silicotungstate 5

Aluminium tungstate,  $\Lambda l_2 W_7 O_{21} 9 \Pi_2 O_7$ , is obtained as a white, flocenlent precipitate by adding a solution of the corresponding sodium tungstate

to an aqueous solution of an aluminum salt 6

Aluminium silicotungstate.— This compound is extremely soluble in water and three hydrates are known. The hydrate  $2\,\mathrm{M}_2\mathrm{O}_3(800_c12\mathrm{W}\,\mathrm{O}_3)$ ,  $93\mathrm{H}_2\mathrm{O}$  crystallises in regular octahedra, the hydrate with  $87\mathrm{H}_2\mathrm{O}$  crystallises in rhombohedra (a=c=1=2.6653), and the hydrate with  $60\mathrm{H}_2\mathrm{O}$  forms triclinic crystals (a=b=c=0.8633=1=1.0658),  $a=87^\circ$  14',  $\beta=105^\circ$  11',  $\gamma=91^\circ$  48'). The first two salts are therefore isomorphous with the corresponding gallium salts (see p 148)

Aluminium phosphotungstate, Al<sub>2</sub>O<sub>5</sub>, P<sub>2</sub>O<sub>5</sub>, 18WO<sub>3</sub>, has been described by Kehrmann 8

<sup>&</sup>lt;sup>1</sup> Nilson, Nova Acta Soc Upsala, 1875, [m], 9, No 7; Ber., 1875, 8, 655, Bull. Soc.

<sup>&</sup>lt;sup>1</sup> Nilson, Nona. Acta Soc. Upsala, 1875, [iii], 9, No. 7; Ber., 1875, 8, 655, Bull. Soc. chim., 1875, [ii], 23, 494.

<sup>2</sup> Weber, Fogg. Annalen, 1859, 108, 615, Wohlwill, Annalen, 1860, 114, 180
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<sup>3</sup> Fairric, Quart. J. Chem. Soc., 1852, 4, 301, Elliot and Storer, Proc. Amer. Acad., 1862, 5, 192

<sup>4</sup> Strive, J. prakt. Chem., 1854, 61, 459, Gentele, ibid., 1860, 81, 411; Parmentier, Compt., rend., 1882, 94, 1713

<sup>5</sup> Copaux, Ann. Chim. Phys., 1906, [viii], 7, 118

<sup>6</sup> Lotr, Annalen, 1852, 83, 65, Anthon, J. prakt. Chem., 1836, 8, 399; 1836, 9, 387.

<sup>7</sup> Wyrouboff, Bull. Soc. pinch. Min. 1898, 19, 219, 1905, 28, 237.

<sup>8</sup> Kehrmann, Zeitsch. anorg. Chem. 1892, 1, 121.

### ALUMINIUM AND THE NITROGEN GROUP.

Aluminium nitride, AlN.—This compound, discovered by Mallet, may be prepared by heating finely divided aluminium in nitrogen at 820° to 1000°, powdering the product, and reheating in the gas once or twice. It is produced when aluminium is heated to 700° in ammonia, and is manufactured by heating a mixture of alumina and carbon in a current of nitrogen.—

$$Al_2O_3 + 3C + N_2 = 2AlN + 3CO.$$

In Serpek's process, a mixture of crushed alumina or bauxite and carbon travels down an inclined rotary kiln, drops into a hopper, and is fed into another rotary kiln, a short length of which is maintained at 1500° to 1800°. Producer gas passing up to the kiln supplies the requisite introgen. The carbon monoxide produced is burned and the hot gases blown up the first kiln to preheat the initial charge.

As usually prepared, aluminum intride forms a grey, amorphous solid. It begins to subline, with partial dissociation into its elements, at c. 1900° (!; the vapour condenses to colourless, hieragonal needles—It is decomposed by water, slowly at 0°, rapidly at 100°, aluminum hydroxide and ammonia being produced, consequently it dissolves readily in alkali hydroxides.

Aluminium phosphide, AIP, prepared from a mixture of aluminium powder and red phosphorus by the thermit reaction, is a friable, yellow solid which is decomposed by water with the evolution of phosphine. The phosphides Al, P<sub>n</sub>, Al<sub>2</sub>P<sub>n</sub>, and Al, P have been described by

phosphides Al,P<sub>1</sub>, Al,P<sub>1</sub>, and Al,P have been described <sup>6</sup>

Aluminium arsenide, AlAs, prepared like the phosphide, is a grey, frieble gold deconversed by water with the evolution of an use <sup>7</sup>

friable solid, decomposed by water with the evolution of arsine  $^7$  . Aluminium nitrate,  $\Lambda l(NO_4)$ , 9H,O, is prepared by dissolving aluminium in nitrio acid of density 1.42 and crystallising the solution. It is dimorphous,  $^8$  crystallising in flat, orthorhombic crystals  $(a \cdot b \cdot c \cdot 0.8925 \cdot 1.10202)$  or in monoclinic pusms  $(a \cdot b \cdot c \cdot 1.1310 \cdot 1.1:1.9191, \beta \cdot 131°36')$  isomorphous with the corresponding ferric salt  $^9$ . It melts at  $c \cdot 70'$ , and at 1.0' is completely decomposed, amorphous aluminium hydroxide being left  $^{10}$ . The aqueous solution when heated deposits basic salts  $^{11}$ 

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    Piccini and Foi Zeit., 1898, 22, 236
    Meyeringh, Ber., phys. nat., 1855, [in.], 16, 460; Eakle, Zeitsch. Kryst. Min., 1896, 6 Church and Northe.
    Klauer, Annalen, 1833. Sci. Petrograd, 1913, p. 407
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    Kluss, Annalen, 1888, 240/him., 1895, [iu.], 13, 48, Ordway, Jahresber., 1858, p. 111.
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89 🌁 ALUMINIUM.

At the ordinary temperature nitric acid of density 1.5 transforms the ennea hydrate into the hexahydiate, Al(NO<sub>3</sub>), 6H<sub>2</sub>O, which is then the stable phase. The hexahydrate often separates from its solution in intric acid of density 1 42, but it is then metastable with respect to the ennea hydrate.1 The transformation from hexa- to ennea-hydrate can be made to proceed in two stages, an intermediate hydrate,  $2\Lambda(NO_i)_1$  [511,0 (or 1611,0 ?), being formed.\(^1\).\(^2\) Ditte \(^1\) has described a dihydrate,  $\Lambda(NO_i)_1$  211,0 \(^2\)
Aluminium Phosphates —The anhydrous and hydrated orthor, mota-,

and pyro-phosphates of aluminium are insoluble in water. Anhydrous aluminium orthophosphate, AlPO,, formed by heating sodium aluminate and phosphoric acid in a scaled tube at 250°, forms colourless, hexagonal prisms of specific gravity 2 59.1 The hydrated orthophosphate is obtained as a gelatmous precipitate by mixing sodium phosphate and neutral aluminium sulphate solutions - It is soluble in ammonia, alkali hydroxides, and immeral acids, but insoluble in acetic acid  $^5$ 

When a solution of aluminium phosphate in an acid is precipitated with ammonia, a basic aluminium phosphate is obtained. Several basic aluminum phosphates occur in nature, e.g. the thombic inmetal wavellite, 3Al<sub>2</sub>O<sub>2</sub>2P<sub>2</sub>O<sub>5</sub>12H<sub>2</sub>O<sub>5</sub> and turquoise. Turquoise (or calaite) is a basic, hydrated aluminum, copper, and ferrous phosphate. It is trichine, has a sky-blue to greenish colour, and is used as a gem stone. It occurs in Persia, Nevada, Arizona, California, and Colorado

Basic aluminium arsenite,  $\Lambda l_2 O_3 \Lambda s_2 O_3$ , and aluminium orthoarsenate, AlAsO, have been described?

# ALUMINIUM AND THE CARBON GROUP

Aluminium carbide, Al<sub>1</sub>C<sub>3</sub>, is prepared by heating fragments of aluminium with sugar carbon in a carbon crucible in the electric furnace and rapidly cooling the product. It may also be prepared by heating a mixture of aluminium powder and carbon for twenty minutes in a Perrot furnace or over the blowpipe, or the reaction may be started by inflaming a mixture of aluminium powder and barium peroxide on the surface. Aluminium carbide is also produced by heating aluminium with hexachlor-

Seligman and Williams, Trans. Chem. Soc., 1916, 109, 612.
 Salm-Horstmai, Juhreeber., 1850, p. 301; Jovitschitsch, Monatsh., 1912, 33, 9.
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 Moissan, Compt. cend., 1894, 179, 16; Pring, Trans. Chem. Soc., 1905, 87, 1630; Askenasy, Jarkowsky, and Waniezek, Zeitsch. Elektrochem., 1908, 14, 811.

benzene at 225°.1 According to Pring, the production of the carbide in the electric furnace is apparently due to the reaction

$$6AI + 3CO_{\sim} = AI_1C_3 + AI_2O_{29}$$

but the elements aluminum and carbon can unite directly, the combination in vacuo becoming perceptible at 650, and rapid at 1400.2

Aluminium carbide, purified by washing with a cold, concentrated solution of potassium hydroxide, and then with alcohol and other, forms yellow, rhombohedral crystals of density at 2.36. At temperatures between 400° and 1100° alummium carbide is stable in vacuo, but above 1400° it dissociates at an approciable rate, and at 1800° the change is rapid. Accordingly, at high temperatures alumina is reduced by earbon with the formation of much free aluminum, while at lower temperatures aluminum carbide only is produced 1 At temperatures below 1400' aluminium carbide acts as a reducing agent on metallic oxides, e.g. .-

$$Al_{1}C_{1} + 12CuO = 12Cu + 3CO_{2} + 2Al_{2}O_{3}$$

but at higher temperatures alloys are produced, the carbon only being oxidised. With metallic chlorides it yields organometallic compounds 5

Aluminium carbide is slowly but completely decomposed by water, alummium hydroxide and methane being produced - 6

$$\Lambda I_4 C_3 + 12 I I_2 O = 4 \Lambda I (O I I)_3 + 3 C I I_1$$

Aluminium carbonate.—The composition of the precipitate obtained from solutions of an aluminium salt and an alkali carbonate varies with the concentrations of the reagents, nature of the aluminium salt, temperature of precipitation, etc. The normal carbonate is never obtained, the precipitates being composed largely of aluminium hydroxide, with smaller quantities of aluminium carbonate, basic aluminium salts, and alkali carbonate?

Aluminium thiocyanate, Al(CNS), is a soluble salt, used in dyeing 8 When solutions of the requisite alkali thiocyanates are evaporated at the ordinary temperature, double salts of the type M3Al(CNS), 111,0 (where M is K, Na, or NH4) are obtained as colourless, deliquescent crystals 9

Matignon, Ann. Chim. Phys., 1908, [vin]. 13, 276, Compt. rend., 1907, 145, 676, Weston and Ellis, Trans. Faraday No., 1908, 4, 60, of Franck, Bull. Soc. chim., 1894, [iii.], 11, 439; Guntz and Masson, that, 1897, [iii.], 17, 209, Dufan, Compt. rend., 1900,

<sup>[</sup>iii.], II, 439; Guntz and Masson, thad, 1897, [iii.], I7, 209; Dufan, Compt. rend., 1900, 131, 541.

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<sup>Piccini and Fo. Zeid., 1898, 19, 707, Gawalowski, Chem. Zentr., 1908, 1-640, Jourdain, 7-Meyeningh, Ber., ph/9602.
Church and Northe. Soc. Chem. Ind., 1883, 2, 589; Hauff, Ber., 1888, 21, 327;
Klauer, Annalen, 183. Set. Voly. J., 1882, 245, 306, G. Stein, ibid., 1883, 250, 36.
(bid., 1838, 44, 472; Proc. Let. 185.900, 33, 1111. Vol alkali aluminium ferrocyaniles, see 10-Kluss, Annalen, 1888, 240/kim., \$, 1353.</sup> 

ALUMINIUM. 91

Aluminium oxalate, Al2(C2O1)3.112O, is precipitated when aqueous solutions of an aluminium salt and an alkali oxalate are mixed soluble in oxalic acid.

When alminimm hydroxide is dissolved in hot, concentrated aqueous solutions of the alkah acid oxalates and the solutions allowed to cool, crystalline double or complex oxidates are obtained of the type  $M_3^1 M(C_2O_1)_3 \times M_2O_4$ The salts are very similar to the corresponding chromic, ferric, and cobaltic double oxalates, and salts having the same number of molecules of water of crystallisation are isomorphons. The salts Ma, Min, (C2O1) p3H2O, for instance, are all monoclinic -

```
K Al – salt 0 9994
Rb Al , 10188
NH, Al ,, 0 9971
                        0 3915 , 92 26
TI Al ", 0 9730
                     1 0 4128 , 93 39
       ., 1 0060
                    1 1 3989 94' 0'
K Ci
Rb Cr. ., 1 0221
NII<sub>1</sub> Cr. ., 0 9830
                        0 3963 92 25
                        -0.3870, 95-18'
K Fe , 0 9918
Rb Fe , 1 0106
                        0.3896 , 94 15
                    1
                        0 4001 , 95 12
                    1 0 3932 , 92 15'
1 0 4115 , 91' 5'
NH<sub>1</sub> Fe ., 0 9959
Tl Fe , 0 9601
NII<sub>1</sub>.Co , 1 0017 1 0 3929 , 92 44
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The following are also isomorphous groups of salts -1

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\begin{array}{ll} \operatorname{Na}_{3}(\operatorname{NH}_{1})_{3}\operatorname{M}_{2}(\operatorname{C}_{2}\operatorname{O}_{1})_{8}\operatorname{7H}_{2}\operatorname{O} & \star \\ \operatorname{Na}_{4}\operatorname{Rb}_{4}^{(1)}_{2}(\operatorname{C}_{2}^{(1)}_{0}_{1}_{0}^{1}\operatorname{7H}_{2}\operatorname{O} \\ \operatorname{Na}_{4}(\operatorname{NH}_{1})_{4}^{(1)}_{2}(\operatorname{C}_{2}\operatorname{O}_{1})_{6}^{1}\operatorname{7H}_{2}\operatorname{O} \\ \operatorname{Na}_{4}(\operatorname{NH}_{1})_{4}\operatorname{Fe}_{2}(\operatorname{C}_{2}\operatorname{O}_{1})_{6}^{1}\operatorname{7H}_{2}\operatorname{O} \\ \operatorname{Na}_{4}(\operatorname{NH}_{1})_{4}\operatorname{Fe}_{2}(\operatorname{C}_{2}\operatorname{O}_{1})_{6}^{1}\operatorname{7H}_{2}\operatorname{O} \end{array}
  Na, Al(C,O1), 511,0
   L<sub>13</sub>Al(C<sub>2</sub>O<sub>4</sub>)<sub>3</sub>6H<sub>2</sub>O
Na<sub>3</sub>C<sub>1</sub>(C<sub>2</sub>O<sub>1</sub>)<sub>3</sub> 5H<sub>2</sub>O
Na<sub>3</sub>Fe(C<sub>2</sub>O<sub>1</sub>)<sub>3</sub> 5H<sub>2</sub>O
Na<sub>3</sub>Co(C<sub>2</sub>O<sub>1</sub>)<sub>3</sub> 5H<sub>2</sub>O
   Li<sub>3</sub>Ci<sub>3</sub>(C<sub>2</sub>O<sub>1</sub>), 6H<sub>2</sub>O
Li<sub>3</sub>Co<sub>2</sub>(C<sub>2</sub>O<sub>1</sub>), 6H<sub>2</sub>O
  (triclinic)
                                  (monoclinic)
  (monochme)
```

Aluminium alkyls. These compounds may be prepared by heating mercury alkyls with aliminium in a scaled tube at 100 They are colour-

less haunds which are decomposed by water <sup>2</sup>

Aluminium methyl, Al(CII<sub>1</sub>), freezes at c 0° and boils at 140°.

Aluminium ethyl, Al(C<sub>2</sub>II<sub>3</sub>), boils at 191′ Aluminium propyl boils at 248° to 252′ Aluminium isobutyl and aluminium isoamyl are also known.

At temperatures only slightly higher than their boiling points, aluminium methyl and aliminium cthyl consist almost exclusively of double molecules  $Al_2(CH_1)_6$  and  $Al_2(C_2H_3)_6$ . The vapour densities diminish rapidly with 1180 of temperature, but it is not possible to state definitely to what extent this diminution is to be attributed to the dissociation Al<sub>2</sub>(Alk)<sub>0</sub> = 2Al(Alk)<sub>p</sub> since it appears probable that decomposition also commences. In ethylene

See Wyrouboff, Bull. Sov. franç. Min., 1900. 23, 65, Copaux. Ann. Chem. Phys., 1905. [vin.], 6 563, Stortenbeker, Ko. trav. chem., 1913. 63, 121, Rosenheim and Colin, Zeitzh. anorg. Chem., 1896, 11, 175, Rosenheim and Platsch, chel., 1899, 21, 1, Lowenstein, 1909, 63, 121. Early references to aluminum oxalates are. Hoese, Compt. ro. ad., 1815, 21, 1116. Mathieu-Plessy, Ibid., 1883, 97, 1033, Collin, Ber., 1870, 2, 315.
 Buckton and Odling, Proc. Roy. Soc., 1865, 14, 19, Calours, Ann. Chem. Phys., 1860, [in.], 58, 5; Hallwachs and Schufarik, Annalen, 1859, 109, 296.

dibromide solution, aluminium ethyl has a molecular weight in harmony with the double formula.1

Aluminium acetylacetonate, [(CH<sub>3</sub> CO)<sub>2</sub>CH]<sub>3</sub>Al, is produced slowly by the action of aluminium hydroxide and rapidly by the action of anhydrous aluminium chloride on acetylacetone. It is best prepared by adding acetylacetone and ammonia to an aqueous solution of aluminium chloride, 2 it separates from its alcoholic solution in white monoclinic (a:b:c=1 901 1: 1.361,  $\beta = 81^{\circ}$  6') 3 crystals isomorphous with the corresponding ruthenium salt,4 melts at 194°, and boils at 315°. At 360° the vapour density is 11.24 (air = 1), the simple formula given above corresponding to the value 11.18 5 The molecular weight also corresponds with the simple formula in solution in benzene and carbon disulphate 6

Aluminium acctylacetonate separates from chloroform solution with two molecules of chloroform of crystallisation. It is insoluble in water and does not combine with aminonia.

Aluminium silicide.—Aluminium and silicon are completely miscible, but no definite compounds are known

The cutectic temperature is 578.7

Aluminium silicates.—The silicate Al<sub>2</sub>SiO<sub>5</sub>, i.e. Al<sub>2</sub>O<sub>4</sub> SiO<sub>2</sub>, occurs in nature as three distinct numerals · cyanite, which crystallises in the furchine system  $(a:b\cdot c=0.899\ 1\cdot0.709,\ a=90^{\circ}\ 5.5',\ \beta\ 101^{\circ}\ 2',\ \gamma\ 105^{\circ}\ 44.5'),$  and alusite, which occurs in rhombic crystals  $(a\ b\ c\ 0.986.1\ 0.703)$ , and sillimanite, which is also rhombic (a  $b \cdot c = 0.9696 \cdot 1 \cdot 0.7046$ ).8 The only one of these forms of the silicate stable above 1300° is sillimanite.

Thermal investigation of the system silica-alumnia shows the existence of only one aluminium silicate, viz Al<sub>2</sub>O<sub>3</sub> SiO<sub>2</sub>, which separates from a fused mixture in crystals of sillimanite, melting at  $1816^\circ$  (fig. 15)  $^0$ 

Aluminium enters into the composition of numerous complex silicates, many of which are important rock-forming minerals. Only a very buef description of the more important of these can be given here.

The felspars are the most important of all rock-forming minerals The chief felspars are orthoclase or potash felspar (monoclinic,  $a \cdot b = c = 0.658.1$ . 0555;  $\beta$  -63° 57') albite or soda felspar (trichine, a b c - 06335 1: 05577, a 91° 3',  $\beta$  - 116° 29',  $\gamma$  -88° 9'), anorthite on time felspar (trichine, a: b: c - 06317 · 1 · 05501; a = 93° 13',  $\beta$  = 115° 55',  $\gamma$  = 91° 12'), and the plagioclases, of which the minorals oligoclase, andesite, labradorite, and The chemical formulæ are KAlSi,Os for orthoclase, bytownite are examples NaAlSi, O8 for albite, and CaAl, Si, O8 for anorthite. As originally supposed by Tschermak in 1864, the plagioclases are isomorphous mixtures of albite and anorthite, 10 Anorthite melts at 1550°.11

Buckton and Odling, loc. cit.; Quincke, Ber., 1889, 22, 551. Louise and Roux, Compt. rend., 1888, 106, 73, 602; 107, 600, V. Meyer, Ber., 1888, 21, 701; Ostwald, Zeitsch. physikal. Chem., 1889, 3, 47.
 Jaeger, Rec trav. chim., 1914, 33, 342.
 Barbieri, Alti R. Accud. Linest, 1914, [v.], 23, i, 334.
 Combes, Compt. rend., 1887, 105, 870, 1889, 108, 405; Bull. Soc. chim., 1889, [iii.],

<sup>Combes, Compt. rend., 1887, 105, 870, 1889, 108, 405; Butt. Soc. chim., 1889, [iii.],
1, 845.
Urbain and Debierne, Compt. rend., 1899, 129, 302; W. Biltz, loc. cit.
Fraenkol, Zeitsch. anorg. Chem., 1908, 58, 154; C. E. Roberts, Trans. Chem. Soc., 1914, 105, 1383.
Shopherd, Rankin, and Wright, Amer. J. Sci., 1909, [iv.], 28, 293, Ballo and Dittler, Zeitsch. anorg. Chem., 1912, 76, 39, Ettal, tbid., 1914, 88, 173.
Day and E. T. Allon, Amer. J. Sci., 1905, [iv.], 19, 93; Dittler, Centr. Min., 1909, 663; Tach. Min. Mitt., 1910, 29, 273.
Day and Sosman, Amer. J. Sci., 1911, [iv.], 37, 341.</sup> 

Orthoclase is mined in large quantities for use in the manufacture of porcelain. Numerous attempts have been made to utilise this mineral as a commercial source of potash. The Ceylon "moonstone," which is used as a gem, consists of orthoclase having a pale blue, pearly opalescence

The micas are numerals of very common occurrence as constituents of rocks. Chemically, they are silicates of aluminum and either an alkali or iron and magnesium, and in addition they contain fluorine and water of constitution. They are monoclinic, but pseudohexagonal, and possess a

highly perfect basal cleavage, which gives white mica or muscoute its commercial importance. Other micas are biotite or dark mica and lepidolite or lithia mica.

The garnets are a well-defined series of minerals of the general formula R "R." (SiO<sub>1</sub>), where R" is Ca, Mg, Fe, or Mn, and R" is Al, Fe, or Cr. The following varieties are distinguished: grossular (Ca,Al), pyrope (Mg,Al), spessarite (Mn,Al), almandine (Fe,Al), uvaroute (Ca,Cr), and and addie (Ca,Fe). The gainets are cube minerals, distinguished by their dodecahedral or icositetrahedral form, greasy lustre, strong refraction, and ready fusibility. Pyrope constitutes the common red gainet used in jewellery.

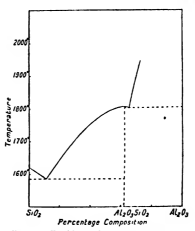


Fig. 15.—Equilibrium diagram for the system alumina - silica

The zeolites are hydrated silicates of aluminium and the alkali metals (and generally of calcium as well). The following immerals are among those classed as zeolites —

The water is very loosely held, and they intumesce before the blowpipe. With loss of water the zeolites lose their transparency, but many of them, after dehydration, can re-absorb the amount of water they originally contained, thereby regaining their transparency and original optical properties. Further, dehydrated zeolites absorb ammoma, hydrogen sulplinde, alcohol, and many other vapours. The nature of these interesting innerals has therefore been the subject of numerous researches <sup>2</sup>

For the literature, see Hart, J. Ind. Eng. Chem., 1912, 4, 827, also Cushman and Coggeshall, ibid., 1912, 4, 821, 1915, 7, 145, Foote and Scholes, ibid., 1912, 4, 377.
 Modern references are as follows. F. W. Clarke, Proc. Washington Acad. Sci., 1905, 257; F. W. Clarke and Steiger, Bull. Geol. U.S. Survey, 1905, No. 262, Ferro, Atts. R. Accad. Linces, 1905, 14, 11. 140. Zambonin, Mem. R. Accad. Linces, 1905, 5, 344; 1906, 6, 102, Rend. Ac., Sci. Fis. Mat. Nappl., 1908, [iii], 14, 148, Zeitsch. Kryst. Min., 1911, 49, 73, Thugutt, Centr. Min., 1909, 67; Grandjean, Bull. Soc. franc. Min., 1910, 33, 5.

Leucite, KAl(SiO<sub>3</sub>), occurs in lavas. It is pseudocubic, crystallising in reositetrahedral forms, and becomes truly cubic at 433°. Nephelite, (K,Na)AlSiO, a hexa mal mineral, is often found in association with leucito. The sodium-alumnium silicate, Na \(\text{ISi}\_2O\_5\), has been prepared artificially. It is known in two forms, the hexagonal form being called soda-nephelite and the triclime form soda-anorthite or carnegierte 1

Spodumene, LiAl(SiO<sub>3</sub>)<sub>2</sub>, is another interesting aluminium silicate. It occurs in large quantities in South Dakota. The green, transparent variety is called haldenite and occurs in North Carolina. The transparent, hlac or violet-coloured variety is known as kunzite, it occurs in California and Madagascar and is a valuable gem-stone. Various lithium-aluminium silicates have been prepared artificially, but they do not agree in physical proporties with any known lithium minerals 3

Topaz,  $Al_2(SiO_1)F_2$  is a fluosilicate of aluminium. It is an orthorhombic innieral (a:b:c=0.528-1-0.177), and its hardness (8), high refractive index, and variety of colour render it valuable as a geni-stone, The usual colour is pale yellow, but light brown, dark brown, and pale blue crystals are also found.

Beryl,  ${\rm Al_2Be_1Sl_6O_{18}}$ , is another beautifully crystalline aluminum mueral. It is hexagonal (hohohedral,  $a \in -1.0$  1989), hardness, 7.5, density, 2.7. The pale green crystals known as aquamarine, and the dark green crystals known as emerald, are familiar as precious stones. Beryl has been synthesised by Hautefeuille and Perrey.1

Tourmaline is a borosilicate of aluminium, alkali metals, iron, and magnesium, containing water of constitution and a little fluorine. It may be regarded as derived from  $\Pi_a\Lambda I_1B_2(OH)_aS_{I_1}O_{I_0}$  by the replacement of the nine hydrogen atoms by metals <sup>5</sup> It is a trigonal mineral (ditigonal pyramidal,  $a = c = 1 \cdot 0.4174$ ), hardness, 7, density, 3.1 Tourmalme is an important tock-forming inneral, it is interesting as an excellent example of a crystal exhibiting polar symmetry and pyro-electric properties, and as a valuable material for the production of plane-polarised light. The colour of tourmaline varies greatly with its chemical composition, from colourless to red, green, brown, and black. Excellent tourmalme crystals, suitable for use as gem-stones, are found in Ceylon and also in Brazil.

Axinite, H('a,Al2BS1016, is also an aluminium borosilicate. It is a beautifully crystalline triclinic mineral (a.b. c=0 1927, 1=0 1511, a=82° 51′,  $\beta$ =88° 9′,  $\gamma$ =131° 33′)

Sodalite, Na (AlCI) Al (SiO1) as a cubic inineral occurring in volcanic rocks; it crystallises in dodecahedra and usually has a blue colour Hauynite. Na<sub>2</sub>Ca(NaSO<sub>4</sub>Al)Al<sub>2</sub>(SiO<sub>4</sub>), is also a blue, cubic immeral, isomorphous with sodalite, with which it is often associated. Hanyinte is of interest as being one of the very few silicates which contain sulphur

Lapis-lazuli is a beautiful blue substance found in Persia, Afghanistan, Siberia, and Chili, and has been prized for ages as an ornamental stone. Density, 2:38 to 2:42, hardness, 5.5. It occurs embedded in limestone, from

Bowen, Amer. J. Sci., 1912, [iv], 33, 551.
 Hess and Wells, Amer. J. Sci., 1911, [iv], 31, 433.
 Ballo and Dittler, Zeitsch. anory. Chem., 1912, 76, 39, Jacger and Simek, Proc. K. Akad. Welensch. Amsterdam, 1914, 17, 239, 251.
 Hautefemille and Pertey, Ann. Chim. Phys., 1890, [vi], 20, 447.
 See Penfield and Foote, Amer. J. Sci., 1899, [iv], 7, 97, Schaller, Zeitsch. Kryst. Min., 1912, 51, 321, Vennadski, ibid., 1913, 53, 273.

which it is extracted by heating the limestone and then breaking out the lapis lazuli with a hammer Lapis lazuli was formerly supposed to be a simple nuneral, but Brogger and Backstrom have shown it to be a rock, consisting of diopside, amphibole, muscovite, calcite, pyrite, and a blue, cubic mineral to which they gave the name lazurite. Lazurite is a sodium aluminium silicate containing both sulphin and chloring and is regarded as being closely analogous and isomorphous with sodalite and hanyinte. Powdered lazurite constitutes the natural pigment ultramarine

**Kaolinite,**  $\Pi_1 M_2 S_1 O_0$ , is a rare, white immerial. It crystallises in the monoclinic system (a, b, c=0.5748, 1=0.5997,  $\beta$ =96, 197), density, 2-62; hardness, 25 The best crystals are found near Amlweh in the island of Anglescy, they are six sided plates or pyramids with a perfect basal cleavage. The mean refractive index is 1.563, the birchingence (0.004) is much lower than that of mica, which resembles kaolinite in appearance? The water is only expelled from kaolinite at high temperatures (above 330"); hence, when the formula is written in the usual way, Al<sub>2</sub>O<sub>3</sub>2SiO<sub>2</sub>2H<sub>2</sub>O<sub>3</sub>te must be remembered that the water is "water of constitution" and not water of crystallisation. Chemically, kaolimite is best regarded as an acid 3.

The question as to the identity or otherwise of kaolinite and the clay

substance of china clay is discussed later (p. 105)

The silicates described above are naturally occurring substances number of artificial aliminium silicates, lowever, are of considerable industrial importance and may therefore be mentioned. These are the substances propared by Gans and known technically as permutits. Sodium permutit is practically an artificial zeolite of the composition 28iO<sub>2</sub>Al<sub>2</sub>O<sub>4</sub> Na<sub>2</sub>O 6H<sub>2</sub>O. It is prepared by heating together china-clay, sodium carbonate and quartz in the requisite proportions and treating the mass with water. Other substances may be added in the preparation, a mixture of 2.5 to 3 parts of china clay, 5 to 6 of alkali carbonate, and 15 to 24 parts of borax, for instance, has been particularly specified. The permutit is of a granular or leafy, porous character

Sodium permutit, like certain natural zeolites, is able to exchange its sodium for other metals with remarkable case. If, for instance, a solution of calcium or magnesium chloride is allowed to percolate through a column of the permutit, a solution of sodium chloride, free from calcium or magnesium, is obtained. If now, a concentrated solution of sodium chloride is percolated through the permutit, the calcium of magnesium is expelled from it and replaced by sodium. The application of sodium permutit to the softening of boiler water will therefore be obvious without further explanation

The sodium or calcium permutits can readily be made to exchange sodium or calcium for manganese, by treatment with a manganous salt solution. When this product is acted upon by potassium permanganate solution, the latter is reduced, and a manganese potassium permutit produced which is covered with very finely divided higher oxides of manganese. This material is a valuable oxidising agent, and, when exhausted, can readily be regenerated by allowing potassium permanganate to percolate through it, its commercial

Brogger and Backstrom, Zetteh, Kryst Min., 1891, 18, 209.
 Dick, Min. May., 1888, 8, 15, 1998, 15, 124
 See p. 96. On the constitution of kaolinite, see Mellor and Holdcroft, vide infra;
 Pukal, cide vitra., Samoilov, Bull. Mod. Sci. Petrograd, 1914, p. 779.
 See G. Schulze, Zeitsch. physikal Chem., 1914, 89, 168.

uses are for removing iron from, and oxidising the organic matter and bacteria in water. These results are easily brought about by allowing the contaminated water to flow through a layer of the permutit; in sterilising water by this means it may be necessary, in addition, first to add a little permanganate to the water.1

Aluminosilicic acids and aluminosilicates.—It is usual to regard the numerous known silicates, including those described in the preceding section, as the salts of various hypothetical silicic acids, or as isomorphous mixtures of such salts.<sup>2</sup> There are, however, good reasons for supposing that many of the compounds usually classed as aluminium silicates are no more related to the silicates than the ferrocyanides, for instance, are related to the cyanides Numerous reasons can be adduced in favour of the view that the alumina in numerous silicates plays the rôle of an acid and not a base, and that, as was originally proposed by Vernadski, the silicates themselves are the salts of a number of complex aluminosilicic acids, comparable with the phosphomolybdic, silicomolybdic, and silicotingstic acids, etc. For the nature of the evidence, however, the reader must be referred to the literature.

Many silicates are decomposed by the prolonged action of water and carbon dioxide, furnishing colloidal silicic acid or opal. In a similar manner, the aluminosilicates give rise to colloidal aluminosilicic acids or clays. The empirical formulæ of these acids may be tabulated as follows .-

| Alumino-monosilicic acid (allophanic type)     | $\Lambda l_{s}O_{s}S_{1}O_{s}nH_{s}O$          |
|------------------------------------------------|------------------------------------------------|
| Alumno-disilicic acid (kaolinic type)          | A1,O,2SiO,n1I,O                                |
| Alumno-trisiliere acid (natrolitie type)       | $\Lambda 1, O, 3S_1O, nH, O$                   |
| Alumino-tetrasilicie acid (pyrophyllitic type) | ALO, $4S_1O$ , $nH$ , O                        |
| Alumino-pentasiliere acid (chabazitie type)    | . $\Lambda l_2^2 O_n^2 .5 SiO_n^2 H_2^2 O_n^2$ |
| Alumino-hexasilicic acid (felspathic type)     | $A1,O,6S_1O,nH,O$                              |

The constitution of these acids at present can only be conjectured. The following groupings of important compounds are due to Mellor and Holdcroft -4

<sup>1</sup> See Gairs, Mitt. aus. d. K. Prufungsanst. f. Wasserversoryung v. Abvasserbesettiquing zu Berlin, 1907, 11cft 8; Woch. f. Brau. 1907, 24, 270 Chem Zeit. 1907, 31, 375; J. Gasbelsucht., 1907, 50, 1925, Jahlb. K. Preuss. Geol. Landesanst v. Benghad. Berlin, 1908, 26, 179, 27, 63, Chem Ind., 1909, 32, 197, Z. Ver Deut Zuckerind., 1907, p. 206, Centr. Min., 1918, pp. 999, 728, 1914, pp. 273, 299, 365, Shemme, whd., 1914, p. 262, Appelius, Chem Rev. Fett Harz, End., 1909, p. 300, Inding and Becker, Chem Zeit., 1908, 32, 514, 531, Anders, Woch f. Brau., 1911, 28, 78, Kolli, Chem. Zeit., 1911, 35, 1393, 1410, L. H. Harrison, Chem. World, 1912, r. 238, Hamor, J. Ind. Eng. Chem., 1912, 4, 240. Also the following patents. Eng. Pat., 1907, Nos. 8232, 3494; 1909, Nos. 21, 184, 26, 842; 1910, No. 28, 353, D. R.P., 1905, No. 174,097, 1908, No. 220,609, Fr. Pat., 1907, No., 374,525, 1909, Nos. 405,990, 409,006, 1910, No. 423,388; 1914, No. 474,283, U.S. Pat., 1908, No., 306,599, 409,006, 1910, No. 423,388; 1914, No. 474,283, U.S. Pat., 1908, No., 913,655; 1910, Nos. 951,641, 968,887.

See Vol. V. of this series.

See Wartha, Annalen, 1873, 170, 333; Brauns, Die chemische Konstitution der Silvate, 1874, p. 6; Gorgen, Ann. Chem. Phys., 1887, [vi.], 10, 145; Vernadski, Bull. Russ. Ges. Naturf., 1891, p. 1, Zeitsch. Kryst. Min., 1901, 34, 37; Zulkowski, Chem. Ind., 1899, 22, 284; Schainer, Zeitsch. Kryst. Min., 1894, 22, 369; Ulfers, J. prakt. Chem., 1907, [ii.], 76, 143; Morozewicz, Kosmos, 1907, 32, 496; Zeitsch. Kryst. Min., 1910, 48, 523; Pukal, Ber., 1910, 43, 2107; Sprechsaal, 1910, 43, 4409 Mellor and Holdcoft, Trans. Eng. Gerama Soc., 1911, 9, 94, 1912, 10, 169; Mellor, Clay and Pottery Industrus Griffin & Co., 1914; W. Asch and A. Asch, The Silvates in Chemistry and Commerce, trans. by Scarle (Constable, 1913), pp. 3–30, where a full discussiony will be found.

97 ALUMINIUM.

# 1. Alumino-monosilicates-Allophanic Type

Allophane is  $\Lambda l_2 O_3 \, SiO_2 \, 5 H_2 O_3$  Salts of this type are -augite, MgO  $\Lambda l_2 O_4$ . SiO<sub>2</sub>, chlorite, 2MgO Al<sub>2</sub>O<sub>3</sub>,SiO, 2H<sub>2</sub>O

### 2 Alumino disdicates - Kaolinic Type

Various alumino disilicie acids are

.  $\frac{\Lambda I_2 O_3}{\Lambda I_2 O_3} \frac{2 S_1 O_2}{2 S_1 O_2} \frac{3 H_2 O}{5 H_2 O}$ Rectorite . Al<sub>2</sub>O<sub>3</sub> 2SiO<sub>2</sub> H<sub>2</sub>O Halloysite Kaolinite M<sub>2</sub>O<sub>3</sub> 281O<sub>2</sub> 2H<sub>2</sub>O Newtonite

Among the aluminosiheates that may be looked upon as salts of the preceding acids are nephelite,  $K_2O/M_2O_3/2SiO_2$ , and anorthite,  $CaO/M_2O_3/2SiO_2$ .

### 3 Alumino-trisilicates - Natrolitic Type.

Salts are represented by nativitie, Na<sub>2</sub>O Al<sub>2</sub>O<sub>3</sub> 3SiO<sub>2</sub>, gainet, 3CaO Al<sub>2</sub>O<sub>3</sub> 3SiO<sub>2</sub>, scalente, CaO Al<sub>2</sub>O<sub>3</sub> 3SiO<sub>2</sub> 3H<sub>2</sub>O<sub>3</sub> tepidolite, KLiO Al<sub>2</sub>O<sub>4</sub> 3SiO<sub>3</sub>, etc

### 4 Alumino-tetrasilicates-Pyrophyllitic Type

Pyrophyllite is  $\Lambda I_2O_1 48iO_2 4I_2O_8$  Salts are levels,  $K_2O_4 M_2O_4 48iO_2$ ; analytic,  $Na_2O_4 M_2O_1 48iO_2$   $24I_2O_8$ , spodumene,  $La_2O_4 M_2O_3$   $48iO_2$ , glaveophane,  $Na_2O_4 M_2O_3$   $48iO_2$ , etc.

# 5 Alumino-pentasilicates—Chaba itie Type

The group is represented by chaberth, CaO M2O3 58iO2 7H2O, and harmotim, BaO M2O3 58iO2 6H2O

### 6 Alumino hemsilicates - Felspathic Type

This type is represented by orthoclass, K2O Al2O3 68iO2

For further information concerning attempts to derive the constitutions of the aluminium silicates, the reader is referred to the literature 1

# ALLMINIUM AND BORON

Aluminium Borides Two bondes are known, AlB2 and AlB12, and crystals have also been prepared containing boron, aluminium, and carbon.2 All the substances were prepared originally by Wohler and Deville, and regarded as allotropic forms of boron crystallising out from solution in molten aluminum Subsequently they recognised that their quaphitic boron was a boride of aluminum  $AlB_p$  but looked upon the aluminum and carbon in thoir adamantine boron as accidental impurities

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<sup>1</sup> The literature references are given by W. Asch and A. Asch, opus ett., where a full account of the various theories of the constitution of the aluminosilicates will also be found, together with a full discussion of the heade postule theory.

2 The literature is as follows. Wohler and Deville, Ann Chin. Phus., 1858, [iii.], 52, 63, Compl. cend., 1867, 64, 19., Innahn, 1867, 141, 268; Hampe, Annahen, 1876, 183, 75; Joly, Compl. rend., 1883, 97, 456, Biltz, Ber., 1908, 41, 2631, 1910, 43, 297, Kuline, D.R.P., 147,871, Binet du Jassonneix, Ann. Chin. Phys., 1909, [viii.], 17, 200

98 5

Aluminium Boride, AlB, -The following methods of preparation are due to Wohler and Deville: (1.) a mixture of boron oxide and carbon is heated to redness in chlorine, and the gaseous products (boron trichloride and carbon monoxide) passed over aluminium heated to whiteness in a porcelain tube; free aluminium is removed from the product by solution in hydrochloric acid or sodium hydroxide; (11) potassium borofluoride (8 pts ), potassium chloride (9 pts ), sodium chloride (7 pts.), and alumnium (5 pts.) are heated together for half an hour at the melting-point of silver, and the product isolated as before Binet du Jassonneix has produced it readily by heating to 1300°, in a stream of hydrogen, a fragment of aluminium contained in a crucible brasqued with boron.

The boude AlBa is a semi-metallic looking solid, which crystallises in sixsided plates belonging to the monochine system 1. It is said by Wohler and Deville, and also by Binet du Jassonners, to be nearly black, but distinctly copper-coloured by reflected light, Joly describes it as golden-yellow in colour. The boride is unaffected by an at a red heat, dissolves slowly in hot concentrated hydrochloric acid or sodium hydroxide, and rapidly in hot intric acid.

When prepared by the method of Binet du Jassonneix, it is accompanied by a small proportion of brown crystals of octahedral habit. These latter crystals are formed in abundance when the boulde AlB, is heated with an excess of boron at 1400°, or when a mixture of boron and aluminium powders is heated to 1300° m hydrogen. They are heavier than the borde  $\overline{\mathrm{AlB}}_2$  and far more resistant towards nitric acid 2

Aluminium Boride, AlB12 -The true nature of this compound was determined by Hampe Binet du Jassonneix has shown that it is produced, . together with numerous other products, when aluminium is heated with excess of boron in a magnesia crucible in the electric furnace. It is best prepared by the method outlined by Kuhne and examined in detail by H. Biltz. Boron oxide (50 grams), sulphur (75 grams), and grampleted Boron oxide (50 grams), sulphur (75 grams), and granulated aluminum (100 grams) are mixed in a clay crucible, covered with magnesium powder, and ignited at the top to start the reaction When cold, the crucible is broken away, the product treated with water, and the aluminum hydroxide and crystalline needles of alumina riused away The lumps of residue are freed from traces of slag, at first mechanically and then by prolonged washing with water. The residue is treated with warm hydrochloric acid for several days; lighter impurities are then removed by decantation with water. The s residue is boiled with hydrochloric acid, then warmed with hydrofluoric acid for some hours, and finally kept in warm, dilute hydrochloric acid for some days.

Thus prepared, the boride forms lustrous crystals, usually six-sided plates, probably belonging to the rhombic system (a.b.c = 7130.1:7139). The crystals appear black, but in thin layers are dark red by transmitted light. At 18° the density is  $2.551 \pm 0.05$ , the specific heats at  $-40^\circ$ ,  $+77^\circ$ ,  $+177^\circ$ , and  $+233^\circ$  are 0.1915, 0.2737, 0.3378, and 0.3663 respectively. The boride is harder than corundum, but softer than diamond. According to Hampe, it is unaffected by concentrated hydrochloric acid or potassium hydroxide solution, is slowly dissolved by hot sulphuric acid, and more readily

<sup>1</sup> Miller, Proc. Roy Soc., 1866, 15, 11, Phil Mag., 1866, [iv], 31, 397.
2 The boron content varies from 66 9 to 79 6 per lent., AlB<sub>12</sub> requires 83 1 per cent.

Possibly they are mixed crystals of AlB, and AlB<sub>12</sub>. Binet du Jassonnex was unable by
this method to observe the formation of the characteristic black plates of AlB<sub>12</sub>.

Weber, Phil. Mag., 1875, [iv.], 49, 161, 276.6

by hot nitric acid. It is oxidised by molten potassium hydroxide or bisulphate, but not by the nitrate, and is attacked also by heated lead chromate.

It rapidly attacks heated platinum, a very fusible product being formed.

Aluminium borocarbides.—The researches of Wohler and Deville and of Hampe were concerned mainly with the interaction of boron oxide and aluminium in either clay or graphite crucibles. Joly showed that when a graphite crucible was employed, the following products were obtained, the relative quantities varying with the conditions of the experiment: (1.) aluminium boride AlB<sub>2</sub>, (ii) aluminium boride AlB<sub>12</sub>, (iii) yellow crystals containing borou, aluminium, and carbon; and (iv.) borou carbide B<sub>6</sub>C. Hampe's experiments led him to believe in the existence of a definite boro-Later, Biltz obtained crystals of definite composition, carbide Al C.B. corresponding to the formula Al<sub>2</sub>C<sub>2</sub>B<sub>11</sub> The method of preparation was similar to that described for the preparation of the bondo AlB<sub>12</sub>, except that soot (2 grams) was added to the mixture. The product, after freatment with water, concentrated hydrochloric acid, and then the warm, dilute acid for some days, consists of sparkling yellow crystals mixed with a few black crystals of  $\tilde{\Lambda} lB_{12}$ 

The crystals have a density at  $18^{\circ}$  of  $2.590 \pm 006$ , are harder than corondum and softer than diamond. Towards reagents they rescrible the

boride AlB<sub>12</sub>, but are more resistant towards immeral acids

It is difficult to believe that the formulæ given by Hampe and Biltz represent definite chemical compounds. Probably each chemist obtained products of definite composition simply because the conditions of experiment were not sufficiently varied. Crystals of quite different composition have been prepared by Binet du Jassonners, by heating aluminium with excess of boson in a graphite crucible in an electric are furnace. They are yellow, transparent, and occur in six-sided plates, mixed with them are crystals of boron carbide, from which they cannot be completely separated. The crystals are attacked slowly by dilute Lydrochlonic acid, quickly by mitric acid After allowing for incchanically admixed boion carbide, the average composition was Al, 64 per cent.; B, 15 per cent. C, 21 per cent., but different preparations varied considerably in composition <sup>1</sup>

Aluminium borate.—The basic borate, 3Al<sub>2</sub>O, B<sub>2</sub>O, was prepared by Ebelinen by strongly heating a mixture of alumina and borax. It forms colourless, orthorhombic crystals (a b c 0 974 1 0.679) of density 3 0.2 The precipitates obtained by mixing alkali borate solutions with solutions of alummum salts are also basic alummum borates."

Aluminium sodium perborate,  $\Delta l_2 \Delta a_2 B_2 O_9 5 H_2 O_8$ , containing 7 to per cent. of active oxygen, is prepared by mixing cold solutions of sodium aluminate (6 6 pts.) and boric acid (5 pts.) and adding 10 per cent.

hydrogen peroxide solution (30 pts) It is a colourless powder, sparingly soluble in water

<sup>1</sup> It is possible that these borocarbides are mixed crystals of two or more of the following substances. All<sub>2</sub>, AlB<sub>12</sub>, Al<sub>4</sub>C<sub>1</sub>, and B<sub>6</sub>C. The suggestion has been made by Binet du Jassonnen that his crystals are mixed crystals of aluminium bonde AlB<sub>2</sub> and aluminium carbide Al<sub>2</sub>C<sub>3</sub>, but it is quite at variance with his analytical data.

2 Ebelmen, Ann Chim Thys., 1851, [iii], 33, 34, Mallard, Compt. rend., 1887, 105, 1260; cf. Troost and Hautefeuille, ibid., 1872, 75, 1819; Frémy and Feil, ibid., 1877, 85, 1029.

3 Rose, Pogg. Annalen, 1854, 91, 452.

4 D.R. P., 285,050.

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### DETECTION AND ESTIMATION OF ALUMINIUM.

Aluminium salts give no precipitate with hydrogen sulphide in acid or neutral solution. Ammonium hydroxide, sulphide, or carbonate, precipitates aluminium as the white, gelatinous hydroxide, which, like the hydroxides of beryllium, vinc, and chromium, is soluble in cold sodium hydroxide. Alumina, when moistened with a solution of a cobalt salt and heated on charcoal, yields a bright blue, infusible mass A test which will detect I part of aluminium in 10 million parts of water is to add I cc of a 01 per cent solution of alizarm S to 5 c c. of the neutral or acid solution, make just ammoniacal, boil for a few minutes, cool, and acidify with dilute acetic acid. A red colour or

precipitate indicates the presence of aluminium.1

In quantitative analysis, aluminium is precipitated as the hydroxide,2 washed, grited over the blowpipe, and weighed as the sesqui oxide Al<sub>2</sub>O<sub>3</sub> i (a) The precipitation is effected, best in a platinum or porcelain vessel, by adding a slight excess of freshly distilled ammonium hydroxide to a slightly acid solution of an aluminum salt, preferably in the presence of ammonium chloride Alternative methods of precipitation are as follows (b) One to three cubic centimetres of phenylhydrazine are added to the hot, very slightly acid solution, the precipitate being washed with a hot, very dilute solution of phenylhydrazine bisulphite free from excess of sulphinous acid 5 (c) The dilute, neutral chloride solution is treated with excess of sodium thiosulphate and boiled until sulphur dioxide is no longer evolved (d). The slightly acid solution is treated with excess of a mixture of potassium iodide and potassium iodate, a moderate excess of sodium thiosulphate added, and the solution heated on the steam-bath for half an hom? (c) The neutral solution is slightly acidified with hydrochloric acid and boiled with excess of ammonium or sodium nitrite, finally, a little ammonia is added

In methods (b), (c), (d), and (c) precipitation is due to the fact that aqueous solutions of aluminium salts are appreciably hydrolysed, and that when excess of a reagent is added which uses up the free acid present, hydrolysis proceeds to  ${\bf completion} \quad {\bf The \ principle \ of \ method \ } (d), " \ {\bf for \ mistance, \ is \ sufficiently \ indicated}$ by the following equations .-

 $\begin{array}{lll} 2AlCl_{8}+6H_{2}O & == 2Al(O1l)_{3}+6HCl \\ K1O_{8}+5K1+6HCl & = 6KCl+3H_{2}O+3l_{2} \\ 6Na_{2}S_{2}O_{8}+3I_{2} & = 3Na_{2}S_{4}O_{6}+6Nal \end{array}$ 

<sup>1</sup> Atack, J Sow Chem Ind. 1915, 34, 936 For further information on the detection of aluminum, see A. A. Noyes, Play, and Spear, J. Amer. Chem. Nov., 1908, 39, 81; Chem. Nov., 1908, 98. 6, etc. Petit, J. Pharm. Chim., 1914, [vir.] 9, 66 (detection as banium aluminate); Rathgen, Zeitsch. anal. Chem., 1913, 53, 33, and Kratzmann, Pharm. Post, 1914, 47, 101, 109 (iniciochemical detection)

2 For the disturbing influence of fluorides, see Verich, J. Amer. Chem. Sow., 1900, 22, 246; Bloor, sbid., 1907, 29, 1603, Curtinan and Dubin, sbid., 1912, 34, 1185, Humiclisch, Ber., 1907, 40, 1497; Zeitsch. anorg. Chem., 1908, 58, 83, Mile. Cavasquae, Compt. rend., 1914, 158, 948.

3 See Daudt, J. Ind. Eng. Chem., 1915, 7, 847.

4 For full details, see Bluin, J. Amer. Chem. Soc., 1916, 38, 1282

9 Hess and Campbell, J. Amer. Chem. Soc., 1899, 21, 776, Allen, sbid., 1903, 25, 421

6 Chancel, Compt. rend., 1858, 46, 987; cf. Hac. Eighth. Inter. Cony. Appl. Chem., 1912, 1, 205.

7 Stock, Ber., 1900, 33, 518

Wynkoop, J. Amer. Chem. Soc., 1897, 19, 434, Schurm, Chem. Zeit., 1909, 33, 877, 1911, 35, 970.

9 Which may be used as a volumetric method (Moody, Amer. J. Sci., 1905, [iv.], 20, 181; Ivanov, J. Russ. Phys. Chem. Soc., 1914, 46,419; Mille. Kovscharova, sbid., 1915, 47, 616; Osipov, sbid., 1915, 47, 618)

The precipitates obtained by methods (c), (d), and (e) are very much easier to filter than that obtained by method (a).

In the presence of phosphoric acid the aluminium precipitate is partly or wholly aluminium phosphate. The precipitation of aluminium as hydroxide is prevented by the presence of organic hydroxy-compounds such as tartaric and citric acids, sucrose, etc., in the solution.

There is no difficulty in separating aluminium from the alkali metals and metals which form sulphides insoluble in acetic acid or dilute inincial acids. The separation of aluminium from magnesium and the alkaline carth metals may be effected by methods (a), (b), (d), and (c) already given. In method (a) the aminoma should be free from carbonate, or it may be replaced by pure aminonium sulphide free from carbonate and sulphate. When precipitated by aminonium hydroxide or sulphide, aluminium may be associated with beryllium, from, chromium, titanium, uranium, zinc, manganese, nickel, and cobalt, though if the hydroxide be used as precipitant, the bulk of the last four metals may be removed, the separation being often sufficiently precise for the purposes of ordinary qualitative analysis.

for the purposes of ordinary qualitative analysis.

The preceding method (e) serves to separate aluminium from zinc, manganese, inckel, and cobalt, as also do the two following methods (f) The cold, slightly acid solution of the chlorides or intrates containing aminonium salts is shaken with excess of pure precipitated barium carbonate and allowed to stand for some hours. The aluminium is precipitated as hydroxide, in accordance with the principle already stated (p. 100), and needs separation from the excess of barium carbonate. (g) The hot, dilute, aeidified chloride solution is made nearly neutral with aminonium hydroxide or carbonate, a slight excess of aminonium acetate added, and the solution boiled for one or two minutes, when the aluminium is precipitated as basic aluminium acetate. Method (f) is more tedious, but more accurate than (g).

Aluminium is readily separated from iron by precipitation with an excess of pure alkalı hydroxide, in which only the hydroxide of aluminium is soluble Alternative procedures are to precipitate the aluminium from a chloride solution by method (c) above, the non present remaining in solution as a ferrous salt, to reduce the non-to-the ferrous state with ammonium bisulphite and employ the phenylhydrazine method (b), double precipitation being advisable, or to dissolve the mixed aluminium and ferric chlorides in hydrochloric acid of density 1 12 and extract the ferric chloride with ether 1 Aluminium is separated from chromium by adding excess of pure alkalı hydroxide, oxidising the chromic hydroxide to alkalı chromate by adding bromme to, or passing chlorine into the liquid, acidifying with nitric acid, and then precipitating the aluminium by the addition of ammonia.2 A good separation of aluminium and titanium is that proposed by Gooch, in which the dilute solution, slightly acidified with sulphuric acid, is treated with sodium acetate and considerable acetic and and boiled for five minutes. Titamum is precipitated as metatitame acid 3. An alternative procedure is to precipitate the titanium with the ammonium salt of nitrosophenylhydroxylamine (cupferron) in a dilute sulphune acid solution. Aluminium may be - - -

Hanriot, Bull Soc. chim , 189°, [ni ], 7, 161.
 Cf Jakob, Bull. Acad Sci (Taron, 1913, Λ, 56, van Pelt, Bull. Soc. chim. Belg., 1014, 28, 125.

<sup>&</sup>lt;sup>3</sup> Gooch, Proc. Amer. Acad., 1884, 29, 435, Chem News, 1885, 52, 55, 68.

<sup>4</sup> Bellucer and Grassi, Atta R. Acad I inver, 1913, [v.], 22, 1 30. Thornton Amer. J.

Sca., 1914, [v.], 37, 407

separated from zirconium in a similar manner 1 It is separated from uranium by precipitation with excess of aminonium carbonate, the uranium remaining in solution

An excellent method for the separation of aluminium from glucinum, iron, zinc, copper, mercury, and bismuth consists in dissolving the mixed chlorides in concentrated hydrochloric acid, adding rather more than an equal volume of ether, and saturating the mixture at 15° with hydrogen chloride gas. The aluminium separates as the hydrated chloride AlCl, 6H,0, which is filtered on asbestos and washed with a mixture of equal volumes of concentrated hydrochloric acid and ether saturated with hydrogen chloride.2 Aluminium nitrate is insoluble in amyl alcohol, a property which may be utilised in separating aluminium from glucinum.3

Details of the various methods must be sought in the original memoirs or in text-books of quantitative analysis 4

<sup>1</sup> Thornton and Hayden, Amer J Sci., 1914, [1v.], 38, 137, Ferian, Atti ist. Veneto scienze lettere ed arti, 1911, 73, 445.

2 Gooch and Hayden, Amer. J Sci., 1896, [iv.], 2, 416, Zeitsch anorg Chem., 1897, 13, 485; Havens, Amer. J Sci., 1897, [iv.], 4, 111, 1898, [iv.], 6, 15, Zeitsch anorg Chem., 1897, 13, 1898, 16, 15, 1898, 18, 147, ct. Minnig, Amer. J. Sci., 1915, [iv.], 39, 197, 40, 482.

Browning and Kuzuian, Eighth Inter Cong. Appl. Chem., 1912, 1, 87 For another method, see Wunder and Mile. Chéhadzó, Ann. Chim. anal., 1911, 16, 205.

4 For the analysis of commercial aluminium and its allova, see Moissan, Compt. rend., 1894, 119, 12, 1895, 121, 851, 1897, 125, 276, Bull. Soc. chim., 1894, [iu.], 11, 1021, 1897, [iu.], 17, 4, Kohn-Abiest, Compt. rend., 1908, 147, 1293, Handy, J. Amer. Chem. Soc., 1896, 18, 766, Seligman and Willot, J. Inst. Met., 1910, 3, 133; Khix, Chem. Zeit., 1911, 35, 668. von John, and., 1913, 37, 363, Crochialski, Zeitsch annew Chem., 1913, 26, 501, Belavo, Ann. Chim. Applicata, 1911, 1, 101, Lunge, Technical Methods of Chemical Analysis, tians. by Keane (Girney & Jackson, 1911), vol. 2, pt. 1, p. 346. For the estimation of aluminium in silicate and carbonate rocks, see Hillebrand, The Analysis of Scheate and Carbonate Rocks (Pail U.S. Geol. Surrey, 1910, Ko. 122), Hinrichen, Ber., 1907, 40, 1497, Selch, Zeitsch anal Chem., 1915, 54, 395. Mellor, A. Treatise on Quantitative Analysis (Griffin & Co., Ltd., 1913), and Treadwell-Hall, Quantitative Analysis (Wiley, 3rd Edn., 1913), may also be consulted with advantage

On the technical analysis of aluminium salts, see Bayer. Zeitsch. anal. Chem., 1885, 24, 542, 1886, 25, 180, Chem. Zeit., 1887, 11, 53, 1888, 12, 1209, 1890, 14, 736, R. Williams, Chem. News, 1887, 56, 194, Beilstein and Grosset. Chem. Soc. Abstr., 1890, p. 85, A. H. White, J. Amer. (Chem., Soc., 1902, 24, 457; Schmatolla, Ber., 1905, 38, 985, Moody, Zeitsch. anorg. Chem., 1907, 52, 286, Chaig. J. Soc. Chem. Ind., 1911, 30, 184. W. W. Scott, J. Ind. Eng. Chem., 1915, 7, 10

### CHAPTER IV.

### CLAY AND CERAMICS.

CLAYS <sup>1</sup> are the chief earthy deposits utilised in the manufacture of bricks, tiles, pottery, and other ceramic products. The term "clay" is one of popular origin and use, and has been incorporated into scientific terminology in a rather loose manner. Geologically, clays occur chiefly among the sedimentary rocks, although they are decomposition products of certain grantes at d some other igneous rocks. Chemically, clays are impure, complex aluminosilicic acids, the characteristics of a clay being dependent on the particular acid present, on its state of hydration, and on the impurities associated with it. Popularly, clays are recognised by the fact that either in a natural state of hidrantic with a sintable quantity of water they become plastic, although different clays vary enormously in this respect, and when heated to redness they become converted into hard, stone-like products

The raw materials from which most clays have been produced are the various felspars, particularly the alkali felspars orthoclase and improcline, and their decomposition products, these immerals were originally present in igneous rocks such as grainte, pegmatite, trachyte, gneiss, etc. Some clays, however, have been derived from rocks containing little or no felspars, e.g. septentine and some gabbros, by the decomposition of augite, homblende, and other aluminosilicate minerals.

The decomposition of the various rock-forming innerals with the production of clay has m most cases been brought about by epigenic or surface actions, or, as is usually said, by "weathering". The rocks expand under the influence of the heat of the sim and contract again at hight, minute cracks being thereby produced. The crystalline structure of the rocks aids the splitting in certain directions, and the disintegration is accelerated by plant roots forcing their way into the eracks and by rain water percolating in and freezing in cold weather. The silicates present in the rocks thus become more and more exposed to the action of surface waters. It has generally

¹ For further information on clays the following works may be consulted · Scarle, British Clans, Shales, and Sainds (C Griffin & Co, Ltd, 1912), Ries, (lays, their Occurrence, Properties, and Uses (Wiley & Sons, 2nd ed., 1909), Searle, The Natural History of Clay (Cambridge University Press, 1912), Searle, The Clayworker's Handbook (C. Griffin & Co., Ltd., 2nd ed., 1911); Nettill, lie Non-inctalize Minerals (Wiley & Sons, 2nd ed., 1910), Howe, A Handbook to the Collection of Kaulin, China Clay, and China Stone in the Museum of Practical Geology (London, 1914); Le Chatelier, La silve et les silveates (Paris, 1914); Morsan, Traité de chinne minérale (Paris, 1905), vol. 1v., article · Céranique °; Thorpe, A Dutionary of Applied Chamistry (Longmans & Co, 1912-13), vols, 11. and iv.; Seger, Gesanwelle Schriften (Berlin, 1896, and American translation, Easton, 1902); and, Mellor, Clay and Pottery Industries, being the Calecter Papers from the County Pottery Laboratory, Staffordshire, vol. 1. (C. Gyllin & Co, Ltd., 1914)

been supposed, following Forschammer, that the carbonic acid held in solution by these waters has been the active agent in effecting the decomposition of the silicates There is evidence, however, that the minerals have also succumbed slowly to the hydrolysing action of water upon them, many powdered, natural silicates being sufficiently soluble in water to react alkaline towards phenolphthalem.2 In the case of orthoclase, for example, it has been suggested 8 that the initial change may be represented thus .-

$$K_2Al_2Si_0O_{16} + 2H_2O \Longrightarrow 2KOH + H_2Al_2Si_6O_{16}$$

the alumnosthere acid then undergoing further decomposition, whereby the elements of silica are partly removed and the elements of water added on until, in general, the final product has the composition II, AloSi2O9.4

Some clays are considered by eniment geologists to have been produced by hypogenic actions, ie actions occurring below the surface of the earth. The Cornish and some of the Zettlitz (Bohemia) china-clays have probably been produced in this manner, the decomposition of the original igneous rocks having proceeded mainly from below upwards, effected by the action of aqueous solutions of carbonic or hydrofluoric acid under pressure and at high temperatures '

Clays which are found overlying or in close association with the rocks from which they have been derived are called primary or residual clays " The chief primary clays are the china clays or kaohus, which are only feebly plastic Clays which have been carried away from their place of origin and deposited elsewhere are known as secondary or transported clays. They are "the washings and sweepings of the hills, which Nature has accumulated as her rubbish-heaps in convenient places," and generally they are very plastic. Some clays, after transportation, have been subjected to great pressures m contact with water, whereby they have become hard and laminated in structure and require to be finely ground before they exhibit any plasticity. Such clays are known as clay-shales, as the result of metamorphic change they pass into slates 7

Composition of Kaolins.—Clays which are white and burn to a white mass are known as kaoline. They may be either primary or secondary clays, but all the kaolins found in the United Kingdom appear to be primary. They occur in association with the igneous rocks from which they were derived, and have to be separated by a process of washing. The bulk of Continental and many extra-European kaolins occur as secondary deposits and need no washing, they differ from English china-clay in several important respects and are usually inferior to it. While the term "kaolin" is employed to denote either a deposit of white clay, a partially decomposed clay-forming rock or the clay that may be obtained from it, separate names are desirable

<sup>1</sup> Forschammer, Pogg Annalen, 1835, 35, 354.

2 F W. Clarke, Bull U.S. Geol. Survey, 1900, No. 167.

3 Cameron and Bell, Bull. U.S. Bureau of Soils, 1905, No. 30.

4 Some chemists consider the true formula to be at least six times as large. See W and D. Asch, The Silicades in Chemistry and Commerce, translation by A. B. Scalle (Constable, 1913), or Sealle, Cement, Concrete and Bricks (Constable, 1913) or Sealle, Cement, Concrete and Bricks (Constable, 1913)

5 Daubiée, Ann. Min., 1841, [in.]. 20, 65; Collins, Vin. Mag., 1887, 7, 205. but cf.

4 Hickling, Trans. Inst. Mag. Eng., 1908, 36, 10, who states that "except in appearance" the product obtained by acting upon felspar with hydrofluoire and "has not the remotest presemblance to china-clay".

emblance to china-clay".

The term residual clay, however, is sometimes used in another sense.

Many other kinds of shales and slates are also known.

for these materials, and in England the terms kaolin, china-clay rock,1 and china-clay are respectively used for the purpose.

English china-clay is separated from the associated rock by directing a stream of water upon it. The finely divided clay is swept away and is carried by the water to a series of catch-pits or pools, in which the sand settles to the hottom. The fluid is pumped to a convenient level and is allowed to flow slowly through a series of long shallow troughs or drags in which the mica and other denser particles settle, leaving the clay in suspension. The finest particles, including all the clay, are then allowed to settle in a pit, after which the clear water is run off, the fine white mild is dired and then forms the china-clay of commerce. The separation into fractions is thus effected by a combination of the processes of sedimentation, based upon the fact that coarse particles sink in water more rapidly than fine ones, and of elutriation, based upon the fact that a slow stream of water flowing over a heterogeneous mass such as a clay carries away with it the small particles more readily than it does the larger ones 2

Hickling <sup>3</sup> has made a careful study of the mineralogical composition of English china-clay, with the following results—The coarse sand which first settles out consists largely of quartz, mixed with tourmaline, biotite (brown mica), and muscovite (white mica). In the succeeding liner saids the quantities of quartz and tournaline diminish considerably while the amount of muscovite increases. The coarse mids which follow contain considerable quantities of kaolinite In the finest muds no biotite is found, tourmaline is extremely scarce, and quartz is only present in small quantities, the bulk of the material consists of kaolimite and the remainder is composed chiefly of muscovite. No felspar is found in English china clay. By exceptionally careful washing it is possible to increase the percentage of kaohiite to per cent., but the best commercial samples only contain 90 to 95 per cent.

Most of the other kaolins appear to resemble the English china-clay in composition, although particles of fel-par are found in various Continental kaolins.

Commercial china-clay is a soft, white or nearly white substance that may easily be reduced to a very fine powder. When mixed with twice its weight of water it will pass completely through a No 200 sieve.

Only a very small portion of commercial kaolins or china-clays can be recognised as the crystalline mineral kaolinite by microscopes of ordinary power,4 the bulk of the material which Hickling claims to be kaolinite appearing to consist of amorphous particles. The identity of this "amorphous" material with crystalline kaohinte is not conclusively proved, and the term clayite, proposed by Mellor, is sometimes used to describe it

The very finely divided muscovite present in commercial china clays is

Felspais 2:55-2 75 Kaolinite 2 6 Biotite 2 7-3:1 Quart 2.65 Mu-covite 2 76-3 00 Calcite 27

<sup>1</sup> Also known as carelazite.

<sup>2</sup> It may here be pointed out that the minerals most plentiful in clays do not differ very greatly in their densities, as shown by the following data.

 <sup>&</sup>lt;sup>3</sup> Hickling, Trans Inst Mag Eng, 1908, 36, 10; J Soc. Dyers, 1915, 31, 70; cf.
 <sup>5</sup> W. Johnson and Blake, Amer J. Sci., 1867, [n.], 43, 351
 <sup>6</sup> The particles present in washed china-clay vary from 0 0005 mm. to 0 03 mm. in

Mellor, Trans Eng. Co. Soc., 1964, 8, 23, Collected Papers, vol 1 p 158. The term clayte was applied some years previously to a mineral of an entirely different nature discovered by H. Clay.

· not the primary muscovite of the original igneous rocks,1 but is most probably produced by the decomposition of felspar. The kaolinite, in its turn, appears to be produced from this secondary mica; the two ininerals are crystallographically very similar, and, on account of their well-developed basal cleavage, occur in china-clay mainly in thin plates 2

The analyses of a specimen of kaolinite and a well-washed Devonshire china-clay are given in the table on p. 108. The analyses of numerous commercial china-clays are in good agreement with the view that the clays

are mixtures of kaolinite, muscovite, and quartz Composition of Transported Clays.—The mineralogical composition

of these clays is naturally more complex than that of the kaolins, since they not only contain the various mineral constituents of the primary rocks and their decomposition products, but also numerous other minerals which have become adventitiously mixed with the clays during transportation from their places of origin to their present sites

Many transported clays are obviously heterogeneous, and, when rubbed up with water so as to form a thin "slip" or "slurry," they may be separated into dissimilar fractions by the use of sieves varying in mesh. The finest fraction may then be further separated by sedimentation or by elutriation. The fractions thus obtained, beginning with that containing the largest grains, are known respectively as gravel, fine gravel, coarse sand, fine sand, silt, fine silt, and clay; and a statement of the composition of a clay in terms of these constituents is termed its mechanical analysis. The following table records the mechanical analyses of samples of Berkshire loam which is used as a brick-clay and Staffordshine "blue brick" clay - 3

|                                                             |   |   | Bakshire<br>Loam. | Staffordshire<br>Clay |
|-------------------------------------------------------------|---|---|-------------------|-----------------------|
| Fine gravel (1-3 mm, diam.). Coatse sand (0 2-1 0 mm diam.) |   |   | 0.3<br>4.4        | 0 8 8 6               |
| Fine sand (0 04-0 2 mm. diam)                               |   | : | 34 8              | 7 2                   |
| Silt (0 01-0 04 mm diam )                                   |   | . | 9.6               | ●8.9                  |
| Fine silt (0.002-0.01 mm diam ) Finest silt                 | • |   | 5·0<br>4·8        | 14 0<br>15 0          |
| Clay .                                                      | • | • | 27.8              | 34 1                  |
| Soluble in hydrochloric acid .                              |   | . | 5 8               | 3 6                   |
| Loss on ignition                                            | • | • | 7 5               | 7 8                   |
| Total .                                                     |   |   | 100 0             | 100 0                 |

The gravel, sand, and silt are found to be devoid of the proporties usually associated with clay. That portion of the original clay to which the "clayey

<sup>1</sup> The primary muscovite may be recognised in the coarse muds when china-clay rock is washed

2 These remarks apply particularly to English china clay, in which the kaolinite may be described as a pseudomorph after secondary muscovite

3 Analyses communicated by Mr A. G. Hopk is; for method of analysis, see Hall,

Trans. Chem. Soc., 1904, 85, 950. Some ceramic chemists adopt other diameters but use . the same names for the fractions.

properties are due, consists of particles considerably smaller than those of fine silt. When the gravel, sand, and silt have been removed from a clay, the residue is often spoken of as "clay substance", this expression is not, however, a very happy one, and it is perhaps better to use the term "argillaceous matter" for the purpose The aigillaceous matter is never homogeneous, and in the case of the commoner clays, such as brick-clays, it may contain notable proportions of such substances as finely divided ferric oxide and calcium carbonate (chalk) unless the clay was treated with hydrochloric acid before analysis. The figures given solve to show the small proportion of "time clay" that exists in many of the commoner clay deposits

Microscopic examination usually reveals the presence of particles of felspar or mica (or both), but in comparatively few clays has the presence of the immeral kaolimite been established with certainty 1. Felspar, mica, and some form of silica (quartz) are so frequently present that they may be looked upon as normal constituents. Of the numerous secondary rock-forming mmerals that occur in clays in small quantities perhaps the most indiquitous is rutile 2 (TiO2) The almost invariable occurrence of titannum in clays was noted in 1862 by Rilev , the amount of titanic oxide, however, seldom exceeds 2 per cent, and in many clay analyses it is (unconsciously) included with the alumina

Of the various other nimeral species present in transported clays the most important are limonite, 2Fe<sub>2</sub>O<sub>1</sub>3H<sub>2</sub>O<sub>2</sub>, hæmatite, Fe<sub>2</sub>O<sub>1</sub>, magnetite, Fe<sub>3</sub>O<sub>4</sub>; siderite, FeCO<sub>1</sub>, pyrite and mareasite, FeS<sub>2</sub>, chalk and limestone, CaCO<sub>3</sub>; magnesite, MgCO<sub>1</sub>, dolonite (Ca,Mg)CO<sub>1</sub>, and gypsum, CaSO<sub>1</sub>2H,O A clay that contains more than 4 or 5 per cent of admixed chalk is called a malm or mail, a clay rich in sand is termed a loam. Small amounts of soluble salts, chiefly sodium, potassium and magnesium sulphates, phosphates and chlorides occur in many clays, and carbonaceous matter is seldom absent, although clays differ widely in the amount they contain. Clays also contain hygroscopic moisture or incchanically admixed water, removable at 105°-110° C, while a further amount of water may be obtained from them at a red heat by the decomposition of the aluminosilicic acids present

The methods used for the ultimate chemical analysis of a clay are those regularly employed in the analysis of silicate rocks. Results are usually expressed as percentages of acidic and basic oxides. Analyses of kaolimite and a number of typical English clays are given in the table on page 108. In certain cases it is necessary to determine more precisely the nature of the "loss on ignition," by estimating separately the water, carbon dioxide, carbon present as organic matter, etc. Moreover, the figure shown for "ferric oxide" is usually only a measure of the total iron present, without reference to its nimeralogical source or sources and its state of oxidation

It is difficult and often impossible to determine the quantitative, or even the qualitative, composition of a transported clay in terms of the minerals

See Hickling, loc cit, for a review of the evidence
 Hussak. Sprechaul, 1889, 22, 136, 151, Teall, Min. Mag., 1885-7, 7, 201, Vogt, La

Hussak, Spiechaud, 1889, 22, 136, 151, Teall, Min. Mag., 1885-7, 7, 201, Vogt, La Céromique, 1905, 6, 68
 Riley, Quart. J. Chem. Soc., 1862–15, 311.
 For the analysis of clays, see especially Mellor, A. Treaties on Quantitative Inorganic Analysis (C. Griffin & Co., Ltd. 1913).
 Analyses 1 and 2 by Mellor and Moderoft (Trans. Eng. Cer. Soc., 1911, 9, 94; 1912, 10, 169), analyses 3, 4, 5, and 6 communicated by Mr. A. G. Hopkins. The data refer to samples dired at 110° C. The mechanical analysis of 4 and 5 is given in the table on p. 106.

present; a fundamental difficulty arises in connection with the nature of the "true clay" present. Of what mineral or minerals is it constituted? In the case of the plastic ball-clays some approach to a definite answer may be given. These clays on careful clutriation and sedimentation yield a residue which consists mainly of amorphous material and, though never entirely free from foreign mineral particles, its analysis approximates very closely to that of the crystalline mineral kaolinite. From this fact it has often been inferred that the basis of all clays is an aluminosilicic acid (or hydrated aluminium silicate) of the formula  $H_1\Lambda l_2 Si_2 O_3$  (or  $\Lambda l_2 O_3.2 SiO_2.2 H_2 O$ ), an inference some-

|                                                                                                                                                                                                                                                                               | 1                | 2                  | 3              | 4                   | 5                  | 6                      |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------|--------------------|----------------|---------------------|--------------------|------------------------|
| Constituent Estimated                                                                                                                                                                                                                                                         | Kaolmite         | China-clay         | Ball-clay      | ". Blue Brick" clay | Red-burning Brick- | Fireclay               |
|                                                                                                                                                                                                                                                                               | (Glamorganshire) | (Lee Moor, Devon). | (Dorsetshire). | (Staffordshire).    | clay (Berkshire)   | (Stourbridge, Staffs.) |
| Silica (SiO <sub>2</sub> ) Titanic oxide (TiO <sub>2</sub> ) Alumina (Al <sub>2</sub> O <sub>3</sub> ) Ferric oxide (Fe <sub>2</sub> O <sub>3</sub> ) Magnesia (MigO) Lime ('uO) Potash (K <sub>2</sub> O) Soda (Na <sub>2</sub> O) Loss on ignition (H <sub>2</sub> O, etc.) | 15 04            | 47·10              | 50.51          | 55 96               | 70:07              | 62·88                  |
|                                                                                                                                                                                                                                                                               | mil              | 0 13               | 0 81           | 0 91                | trace              | 0 73                   |
|                                                                                                                                                                                                                                                                               | 38 58            | 39 42              | 34 22          | 19:61               | 13:45              | 25·26                  |
|                                                                                                                                                                                                                                                                               | 0 14             | 0 23               | 1 74           | 9 51                | 6 09               | 1 71                   |
|                                                                                                                                                                                                                                                                               | 0 01             | 0 24               | 0 02           | 1:45                | 1 05               | 0 04                   |
|                                                                                                                                                                                                                                                                               | 0 38             | 0 31               | 0 43           | 0 65                | 0:66               | 0 33                   |
|                                                                                                                                                                                                                                                                               | 0 18             | 0·16               | 0 81           | 1 28                | 1 38               | 0 19                   |
|                                                                                                                                                                                                                                                                               | 0 13             | 0 08               | 1.65           | 1 23                | 1 39               | 0 28                   |
|                                                                                                                                                                                                                                                                               | 15 94            | 12 24              | 9 59           | 6 27                | 4 77               | 8 87                   |
| Total .                                                                                                                                                                                                                                                                       | 100 40           | 9 <b>9</b> 91      | 99 81          | 99 87               | 99 16              | 100 29                 |

times expressed by saying that clays are composed of true clay or kaolimite associated with various other ininerals. It is, however, very undesnable that the name kaolimite should be employed to denote anything but the crystalline mineral, and it is at least premature, and most probably erioneous, to conclude that only one aluminosilicio acid exists in clays. A number of minerals which are presumably aluminosilicio acids of a somewhat similar nature to kaolinite, e.g. allophane, collyrite, pholerite, nacrite, rectorite, halloysite, newtonite, pyrophyllite, montimorillonite, etc., are known. It is not certain that all these substances (several of which are only known in the amorphous state) are distinct mineral species. several of them may possibly be mixtures containing silicio acid and aluminium hydroxide. The detection of these substances in clays is very difficult, since, with the exception of the china-

<sup>1</sup> This includes 14:44 per cent H<sub>2</sub>O and 0:80 per cent of carbon

2 The literature relating to these substances is very confusing. For example, Le Chatcher

(gives collyrite the formula Al<sub>2</sub>O<sub>3</sub>, 28:10, 6H<sub>2</sub>O and regards allophane as identical with it,

others give allophane the formula Al<sub>2</sub>O<sub>3</sub> 8:10, 5H<sub>2</sub>O, 8 See Doelter, Handbuch der Mineral

chemie, vol. 11. (1914); Ries, opins cit, Searle, British Ulays, Shalis, and Sands (C. Griffin

& Co., Ltd., 1912) W and D Asch, The Silvates in Chemistry and Commerce, transl. by

A. B. Searle (Constable, 1913); Searle, Ciment, Concrete and Bricks (Constable, 1913).

clays and ball clays, the "true clay" present can rarely be isolated in anything like a pure state, from the work of van Bennmelen, however, it appears that collyrite, the only member of the series that is decomposed by hydrochloric acid, is present in numerous clays 1. Le Chatcher 2 has proposed to identify the different aluminosilicic acids by means of certain thermal changes that may be observed when they are heated, and by this method he claims to have established the identity of halloysite and pholerite with the alumino silicic acid present in china-clay.

At the present time it is usually considered that the "time clay present in a transported clay consists of aluminosilicic acid or mixture of such acids, sometimes with free silicie acid and aluminium hydroxide! Although in the case of the ball-clays (and probably numerous other clays) the "true clay" has the same composition as knolimite, it is not correct to consider that it is an amorphous form of that mineral, or that it is identical with elayite (p. 105) It differs from the latter firstly in that its particles are much smaller and do not seem to have been shown to be crystalline, and secondly in that it is extremely plastic. Accordingly, Searle 5 distinguishes the "true clay" of ball-clays and other highly plastic transported clays by the term pelinite

Attempts to determine the inmeralogical composition of clays are often made, based upon the fact that the "true clay" they contain is attacked by concentrated sulphuric acid. When a clay is heated with concentrated sulphuric acid the "true clay" contained in it is decomposed with the production of aluminum sulphate and silicic acid hydrogel, and these may be removed by digesting the mass alternately with sodium hydroxide or carbonate and hydrochloric acid. The loss in weight which the clay undergoes is often regarded as representing the "true clay" or "clay substance" present. The non-plastic residue is then regarded as being a mixture of felspar or mica and quartz. The silica and alumina it contains are determined and the equivalent amounts of felspar or mea and quartz are calculated

An analysis of clay carried out according to the scheme just outlined is called (or rather miscalled) a rational analysis. It is based upon the assumption that only the "true clay" is rendered soluble by sulphuric acid, whereas it is known that mica is always more or less attacked, the extent varying with the nature of the mica, its state of subdivision, the concentration of the acid, and the time of heating. Felspar is also attacked, but not so readily as mica Hence the material decomposed by the sulphune acid

Van Bemnielen, Leitsch-auberg Chem., 1904, 42, 265., 1909, 62, 221., 1910, 66, 322.
 Le Chatcher, Compt. vend., 1887, 104, 1143, 1517. Bull. Soc franç. Men., 1887,

<sup>10, 207</sup> 3 Le Chatelier, La silve et les silvates (Paris, 1914).

<sup>3</sup> Lo Chatcher, La silve et les silveites (Paris, 1914).
4 Clays rich in free aluminium hydroxide are termed laterite days or laterites and are particularly abundant in the tropics. It is supposed that under tropical conditions ordinary clays are slowly decomposed into free silica and alumina (see van Bennielen, loc cit; Lacroix, Comp. rend., 1914, 159, 517. Fermor, Grot May, 1915, 2, 28, 77, 123). It has been suggested that the hearists (P 73) are clays in which the lateriation is very advanced.

Scarle, The Natural History of Clay (Cambridge University Press, 1912), p. 148.

According to Asch and Asch (opus cit) the effect of heating true clay with sulphinric acid is not the production of aluminoum sulphate and silica card but the formation of a romalkably stable soluble aluminoulic anhydride and the separation of some free silica. In the pure primary clays the number of aluminosilicic acids present is limited and the composition of the soluble anhydride is consequently constant, but transported clays have usually derived their "time clay" from several sources so that it probably contains several aluminosilicic acids and the true nature of the mixed soluble anhydrides cannot then be ascertained.

is not merely the "true clay" present. Again, many of the commoner clays contain considerable quantities of chalk, siderite, and hiematite, or notable amounts of colloidal silicid acid or organic matter, which must be determined separately and their influence on the rational analysis allowed for; hence whilst with the purer kaolins the errors may be only small, with common clays this incthod of analysis is of little or no value and is frequently misleading. In fact, many rational analyses of clays recorded in the literature are quite inconsistent with the ultimate analyses of the same clays. Hence, as Searle has pointed out, a consideration of the ultimate chemical analysis and a microscopic examination of a clay is, at present, the most satisfactory method of arriving at an approximate idea of its mineralogical composition.

Physical Properties of Clays.—Most clays are fairly soft solid bodies, particularly when moistened with water, they have a characteristic "argillaceous" odour, which is usually attributed to the presence of organic matter. The true density usually varies between 2 50 and 2 65, but as some clays are more porous than others the apparent density varies very considerably

Freshly dug clays vary very considerably in colour, from nearly white to almost black; most commonly they are yellow, grey, or greyish-brown. The colour is due partly to the organic matter present and partly to the presence of certain innerals, usually compounds of iron (hamatic, limonite, siderite); colour is accordingly no reliable criterion of the purity of a clay or of the colour it will become when burned in a kiln or pottery oven

A very obvious and extremely important property of a moist clay is its plasticity, or ability to change its shape without rupture when submitted to pressure and to retain its new shape when the pressure is removed. Different kinds of clays vary enormously in their degrees of plasticity. No entirely satisfactory method for measuring the plasticity of clay has yet been devised.

A decidedly plastic clay has the power of remaining plastic when other finely divided but non-plastic materials, e.g. said and brick-dust, are incorporated into it. In general, the more plastic a clay the greater the amount of non-plastic material which it can thus "bind," and the more finely divided the added substance the smaller is its effect in reducing the tensile strength of the clay.

Owing to its heterogeneous character a dry clay is porous, i.e. its apparent volume includes the volume of a considerable quantity of air, this may be roughly estimated by determining the quantity of met liquid which it can absorb without changing its volume. If a little water is added in small quantities the clay first becomes granular and then pasty. The addition of a still further quantity of water separates the particles to such an extent that the cohesion is destroyed, the clay "runs," and a fluid clay "shp" or "slurry" is produced

When a clay that has been kneaded into a plastic mass with water is set aside in the open, evaporation of water commences at the surface. As the surface water evaporates it is replemished from within the mass, and as long as there is any water between the clay particles the process of drying is attended by a shrinkage in the volume of the mass. A stage is eventually reached when the solid particles are in direct contact with one another,

<sup>2</sup> Searle, opus cit.

<sup>.. 1</sup> Mellot, Collected Papers, 1914, vol 1 p. 109, and A Treatise on Quantitative Inorganie
.. Analysis, 1913 (C Griffin & Co., Ltd.), chap. xliv., where a critical discussion of the method
of rational analysis will be found.

and the removal of the remaining small amount of water is not attended by any further shrinkage. The amount of shrinkage depends upon the texture of the clay, the relative proportions of plastic and non-plastic materials it contains, the amount of water added, and several other factors, and a shrinkage in volume of 12 to 38 per cent may be considered normal. In the preparation of various ceramic products from clays which exhibit a large airshrinkage, sand or material of a sandy nature is often added to reduce the shrinkage, and an additional advantage is thereby gained as the increased porosity permits the mass to be dried more rapidly and minimises danger from cracking.

Many theories have been advanced to account for the plasticity of wet clay, but not one can be considered as wholly satisfactory. 

In a general way it may be stated that each solid particle present in the clay must be surrounded by a film that is coherent and self-healing? In the opinion of Le Chatcher s the film is composed merely of water, the coherency of the plastic mass being attributed to (i.) the lamellar structure of the clay particles, which can accordingly slip over one another readily like the various units in a pack of cards, and (ii ) the presence of minute air-bubbles in the mass, on account of which innumerable capillary menisci a o produced and the lamellar particles held tightly together by the operation of

It is an undoubted fact that very finely divided 4 kaolinite, inica, glaucophane, calcite, gypsum, and other minerals which readily cleave into plates, acquire a certain plasticity when inixed with the requisite amount of water; yet, while it is possible that Le Chatelier's explanation of plasticity may account satisfactorily for the feeble plasticity of the kaohus, it is improbable that it can be more than a mmor cause of the plasticity of the highly plastic secondary clays, even if it be granted that with a smaller size of clay particle the plasticity would be augmented

It is difficult to connect plasticity with chemical composition, as clays which yield almost identical results upon analysis may differ widely in plasticity, moreover, clays that are inchest in "clay substance" are often less plastic than those which are less pure, so that although many authorities have attributed plasticity to a peculiar structure of the clay molecule, such

an explanation should not be hastily accepted.4

There is considerable evidence for the view that most clays contain colloid matter,' and that the plasticity of a clay is closely connected with the quantity of colloid matter present. Ashley, in a discussion of this subject,6 considers that both morganic and organic colloids may occur in clays. The former include amorphous aluminosihere acids, sihere acid, ferric hydroxide, and occasionally aluminium hydroxide, the latter, which have been produced from vegetable matter, are usually referred to as "humins"

<sup>1</sup> The more important theories have been carefully summarised by Scarle, British Clays,

<sup>1</sup> The more important theories have been carefully summarised by Scarle, British Clays, Shules, and Sands (C. Guilin & Co., Ltd., 1912)

2 Bancrott, J. Physical Chem., 1914, 18, 783

3 Le Chateher, opus cit. (p. 109). See also S. W. Johnson and J. M. Blake, Amer. J. Sci., 1867, [n.], 43, 351.

4 For an attempt to connect the plasticity of clay with the chemical constitution of the clay molecule, see W. and D. Asch, opus cit.

5 First suggested by Way, J. Roy. Agric. Soc., 1850, 11, 313, and Schloesing, Compt. 1end., 1874, 79, 376, 473.

5 Ashley, Bull. U.S. Geol. Survey, 1909, No. 388, where numerous references to the literature may be found. literature may be found.

or "humic acid." It has been suggested by some that the organic colloids are the most important,1 but this cannot be regarded as proved. On the colloid theory of plasticity, a wet, plastic mass of clay is regarded as consisting of innumerable minute solid mineral particles disseminated throughout a gelatmous, colloidal hydrogel or mixture of hydrogels, saturated with

 $\Lambda$  hydrogel such as precipitated silicic acid shrinks very largely in volume on drying, and when dired at a fairly low temperature forms a very hygroscopic mass which can take up a considerable amount of water without appearing wet. In these respects it closely resembles a plastic clay-Plastic clays are in various other respects similar to hydrogels, e.g. they adsorb substances selectively from aqueous solutions. The adsorbed substances may be either metallic salts or organic dyes of a colloidal nature.2 The adsorption of 1 or 2 per cent. of certain organic colloids such as proposed to measure the plasticity of a clay by its power to adsorb methylene blue

The presence of reversible colloids in a plastic clay is suggested by the manner in which the viscosity of clay slips is influenced by the presence of metallic salts 4 For instance, a plastic clay which has been worked up with water into a very viscous mass may be transformed into a mobile liquid by the addition of a little sodium hydroxide or carbonate, and reconverted into a viscous mass by the further addition of a little hydrochloric acid The explanation is probably as follows. The colloids present in clays are negatively charged when in colloidal suspension, and accordingly it is mainly the positive ions of the electrolytes added that are of influence in producing their coagulation. All clays contain soluble salts to some extent, although it may often be but a slight one, and of the ions into which they dissociate the positive, bivalent calcium ion has most effect in preventing the hydrogels present from passing into hydrosols when the clay is stirred up with water The addition of sodium carbonate leads to the removal of the calcium ions and their replacement by univalent sodium ions which, being of lower valency, are less able to prevent the conversion of gel into sol.6 On this view the addition of sodium carbonate little by little would be expected first to facilitate sol formation and thus dimnish the viscosity until the calcium was precipitated, and then, with mereasing concentration of sodium ions, to facilitate the reverse change from sol to gel and so increase the viscosity? This is precisely how the viscosity is observed

<sup>1</sup> See, e.g., Stewart, J. Ind. Eng. Chem., 1913, 5, 421

2 Way, J. Roy. Agr. e. Soc., 1850, 11, 313, Hursch, Tonind. Zeil., 1901, 28, 491, Ries, Traks. Amer. Cr. Soc., 1904, 6, 44; Ashley, loc. eil.

3 Acheson, Trans. Amer. Cer. Soc., 1908, 16, 31, 527, 881, 1025, 1906, 39, 169, 1167, 1184; Bleininger, Trans. Amer. Cer. Soc., 1908, 10, 389; Mellor, Green, and Baugh, Trans. Eng. Cer. Soc., 1908, 6, 161, or Mellon, Collected Papers, vol. p. 88, Back, Trans. Amer. Cer. Soc., 1914, 16, 515; Bleininger, J. Franklin Inst., 1915, 180, 225

3 W. and D. Asch (opus cit.) consider that the addition of small quantities of alkaliconvoits clays into strongly acid salts by replacing only one or two hydrogen atoms in a highly complex molecule. They also consider that the addition of a weak acid replaces the alkali metal by hydrogen and restores the original clay molecule. They find confirmation of their views in the differential behaviour of china-clay, which had been treated with soda, to strong and weak acids respectively. to strong and weak acids respectively.

See Vol. I. p. 84.

F. Foerster, Chem. Ind., 1905, 28, 733.

to vary.1 The great value of sodium carbonate in the production of a fluid slip containing a minimum of water has been known for many years and is utilised in preparing ceramic bodies by the method of slip easting (p. 126).

W. and D Asch claim that their theory of the molecular constitution of clays accounts for all the so-called colloidal properties in a purely chemical manner.

Action of Heat on Clays. - The simplest case is that of the action of heat on china-clay that has been washed free from all but a very small percentage of mica and quartz, i.e a mixture of clayite and kaohinte, the former being in great excess. The decomposition of this mixture by heat has been carefully studied by Mellor and Holdcroft.2 At 100° to 110" C the hygroscopic moisture is driven off, at higher temperatures water is evolved in consequence of the breakdown of the molecule. It is not possible to state any definite temperature below which this decomposition ceases, the rate of decomposition increases with rise of temperatine and is quite perceptible at 500° C, the mass losing weight at the rate of 12 per cent per hom. As the temperature is increased to a little above 500', decomposition of the clay occurs with a decided absorption of heat (about 10 8 Cals per formula-weight in grains of clayite). The evidence is not conclusive on the matter, but Mellor and Holdcroft think it very probable that this heat absorption corresponds with the following chemical change -

$$H_1AI_2Si_2O_9 = AI_2O_3 + 2SiO_2 + 2H_3O_3$$

When the temperature reaches a little more than 800° a marked evolution of heat is observed, owing to some kind of change (polymerisation !) taking place in the alumina (cf. p. 72).4 At temperatures between 1200° and 1300° the silica and alumma commence to reunite at a slow but perceptible rate, for crystals of sillmanite, M.O. SiO. begin to form in the mass. At higher temperatures (above 1700), owing to the fusion of the traces of mica and silica initially present and the silica produced from the decomposition of the clayite, the mass commences to sinter, and it becomes decidedly fluid at 1710' C.º

The thermal changes observed with china-clay at temperatures just above 500° and 800° are also noticed when the pure immeral kaohinte is heated;7

<sup>1</sup> For further information on the colloidal theory of plasticity, see Ashley, loc. cit.; Rohland, Die Tone (Vienna, 1909). Rohland, Lusch, among Chem, 1902, 31, 158; Spreksaul, 1906, 39, 129, 1371, Bowlem Ledich, 1912, 46, 374, Koltond Ledish, 1914, 14, 193, Cushman, J. Amer Chem Soc., 1903, 25, 451, Trans Amer Ce. Soc., 1904, 6, 65; N. B. Davis, blid, 1914, 16, 65, Bull Amer. Inst. Mag. Eng., 1915, p. 301, and special treatises on class.

2 Melloi and Holdcroft, Trans Eng. Cer. Soc., 1911, 9, 91, Mellor, Colloided Papers, 1914, vol. 1, p. 722. See also Le Chatcher, ones att., and Bull. Soc. Lunc. May. 1887, 19, 207.

<sup>&</sup>lt;sup>2</sup> Mellot and Holdcroft, Trans. Eng. (7), Noc., 1911, 9, 91, Melter, Cottoted Papers, 1914, vol. 1, p. 272. See also Le Chatcher, opas cit., and Bull. Noc. frame. Min., 1887, 10, 207; Compt. read., 1887, 104, 1443, 1517, Samonlov, Bull. Acad. Sci. Petrograd, 1914, p. 779.

<sup>3</sup> All that is certainly known is that water is given off in the proportion indicated in the equation, and that the residue is soluble in dilute hydrochloric acid. This is micropatible with the production of free silies, and is regarded by some chemists as indicating the formation of a soluble aliminosilicie anhydride Al<sub>22</sub>Si<sub>2</sub>O<sub>374,2y</sub>, which polymerises or decomposes and becomes insoluble on further heating. See W. and D. Asch, opas cit.

<sup>4</sup> Or in the aliminosilicie anhydride (4) necessiting featings.

Or in the aluminosilicic anhydride (cf. preceding footnote).
Of the aluminosilicic anhydride to decompose.

<sup>&</sup>lt;sup>b</sup> Kanolt, J. Washington Acad Sci., 1912, 2, 337, Tech. Papers, Bur. Stand., 1912,

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Mellor and Holdcroft, Trans Eng Cer. Soc., 1912, 10, 169, Mellor, Collected Papers, 1914, vol. 1. p. 297.

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this is a strong argument in favour of the view that clayite and kaolinite are the same substance in different degrees of subdivision or different physical conditions

Little is known of the action of heat upon the other aluminosilicic acids which are thought to exist in clays, but it is at least very probable that at temperatures above 500° or thereabouts they behave his kaolimite.

Ordinary clays are heterogeneous, and hence cannot be said to have melting-points in the true sense of the term. Of the variously finely divided minerals present in them, some, e g. the felspars and micas, have definite meltingpoints, while others, eg chalk, dolomite, siderite, pyrites, gypsum, etc., when heated, undergo chemical decomposition at temperatures below the meltingpoints of the former, with the production of substances of very high melting-point (lime, magnesia, ferrous oxide, etc.). These substances, however, are basic in chemical character, and react with silica, micas, and felspais at temperatures of about 900' to 1000', i.e. considerably below the melting points of these substances,3 with the production of molten silicate mixtures. As the heating is continued and the temperature rises, the felspars and micas, still unchanged, commence to melt. The felspars, micas, lime, magnesia, etc., in a clay are accordingly said to be the fluxes present. It is a curious fact that, except in the presence of lime, ferric oxide does not appear to act as a flux but rather as an infusible material, ferious oxide, however, is a powerful flux. Titanic oxide is an acidic oxide which, like silica, acts as a flux in refractory clays.

The fusibility of a clay depends upon the nature and amount of the fluxes present. Bearing in mind that feiric oxide may or may not act as a flux, the amount of fluxes present may be seen from the ultimate analysis, the sum of the alkalies, lune, magnesia, "forme oxide," and titanic oxide is often spoken of as the amount of fluxing impurities contained in the clay. Concerning the nature of the fluxes present, it may be said that the alkalies are contained almost entirely in the micas and felspars, which numerals also account for at least part, and sometimes practically all, of the lime and magnesia present. The presence of above 5 per cent. of line is, however, an almost certain

indication that free calcium carbonate is present in the clay

The action of heat on a natural clay is such that at a certain temperature, which may be as low as about 900° C or several hundred degrees higher according to the nature of the clay, molten material commences to form in it This fluid, which is very viscous, acts as a kind of "cement," binding the still unfused particles together, on cooling, the "cement" sets to an amorphous "glass," instead of crystallising, and, unless the proportion of fluxes is too small or the time of heating too short, a hard porous mass having considerable crushing strength is obtained

<sup>1</sup> See, however, Le Chatcher, loc ad

<sup>&</sup>lt;sup>1</sup> See, however, Le Chatchet, loc cd
<sup>2</sup> E g, the dissociation-pressure of calcium carbonate reaches the value of one atmosphere at 898° C. (Johnston, J. Amer. Chem. Soc., 1910, 32, 933).

<sup>3</sup> The melting-points of the alkali felspais albite and orthoclase are not known with much accuracy, but are rather lower than 1200° C,; Bowen gives the value 1100±10° C, for albite. The melting-points of the plagnoclase felspais rise steadily with increase of calcium content from c. 1100° C, the melting point of albite, to 1550° C, the melting-point of anorthite (Day and Allen, Amer. J. Sci., 1905, [v.], 19, 93; Day and Sosman, ibid, 1911, [v.], 31, 341, Bowen, ibid., 1913, [v.], 35, 577). See also Joly, Proc. Roy. Irish Acad., 1891, [iii], 2, 33; Cusack, ibid., 1896, [iii], 4, 339; Doelter, Tsch. Min. Mitt., 1901, 20, 210, 1902, 21, 23. White nuca or nuscovite is not such a readily fusible flux as an alkali felspai; see R. T. Stull, Trans. Amer. Cer. Soc., 1902, 4, 255.

If the period of heating is prolonged, more and more of the fluxes enter into the fluid state. The surface tension between fluid and solid causes the solid particles to be drawn closer and closer together, and this, with the flow of fluid into the pores, diminishes the porosity of the heated clay. At the same time the fluid exerts a solvent action upon the solid particles. The entire process is spoken of as the vitrification of the clay. Tho speed at which the vitrification proceeds at a given temperature depends upon the composition of the clay and on its textine, i.e. the sizes of the various kinds of particles present: quartz, for instance, is more readily dissolved by the molten fluxes when very finely divided than when it is present in coarse grains. By raising the temperature, the rate of vitrification is accelerated

When the period of heating of the clay is sufficiently prolonged at an assigned temperature, it may happen that the vitrification is carried so far that the mass is no longer capable of retaining its shape, i.e. it softens. The more readily a clay softens the less refractory it is said to be. The softening of the mass is determined by the ratio of the fused to the unfused material present, i.e. by the extent to which vitrification has proceeded, and by the viscosity of the fused material. As the vitrification is caused by slow reactions in heterogeneous systems, the temperature of softening is not a measure of the refractoriness of a clay without reference to the manner in which it is heated, for the temperature of softening will vary with the rate of use in the temperature of the clay, with the chemical changes undergone by the clay, with the use of an oxidising or a reducing atmosphere, and with the pressure, since a viscous fluid which is meapable of flowing under its own weight may be caused to flow by subjecting it to sufficient pressure.

When a clay is heated to a temperature at which vitrification will proceed, and maintained at that temperature, it does not follow that vitrification will continue until the mass softens. If the percentage of fluxes is low, the fluxes soon become saturated with the retractory particles and form a solution which is practically infusible at the given temperature. The temperature must then be raised before the fluxes can again resume their work of dissolving the refractory particles while drawing them closer together. On the other hand, the proportion of fluxes may be such that, although the mass does not soften, vitrification continues at the given temperature until the porosity of the mass has been reduced practically to zero and a stoneware mass, impervious to water, has been produced. The clay is then said to have been completely vitrified, and clays which behave in this manner are called vitrifiable or stoneware clays.

The extent to which vitrification proceeds increases with rise of temperature, so that a clay which completely vitrifies at a given temperature may not do so at another and higher temperature without softening and losing its shape. The temperature interval between the point of incipient vitrification and the softening point is spoken of as the "range of vitrification." Since clayware loses its commercial value when it softens and loses its shape, it will be clear that the production of stoneware from vitrifiable clays is only safe with clays having a long range of vitrification, if the range is short, disastrous results may easily occur as the result of slight overheating. It has been found that calcareous clays are those which have the shortest range of vitrification. In this respect the fluxing action of lime is in marked contrast with that of magnesia.<sup>1</sup>

<sup>&</sup>lt;sup>1</sup> Mackler, Tonind Zeit, 1904, 26, 706; Hottinger, Trans. Amer. Cer. Soc., 1903, 5, 180; Barringer, ibid., 1904, 6, 86.

The softening point of a clay is usually determined by forming a test portion into a small triangular pyramid about five or six times as high as the length of its base, drying it, mounting it vertically in a suitable furnace, and then raising its temperature until the pyramid, or "cone" as it is usually called, bends over and its apex reaches the level of the base. According to Mellor, if the temperature is then to C and the cone commenced to bend at  $t_1$ ° C., the interval  $t_1$  to  $t_2$  represents the softening range and the mean temperature  $(t_1 \times t_2)/2$  may be taken as the softening point throughout the heating a reducing atmosphere should be avoided, and the temperature should be raised at the (arbitrary) rate of 10° C per minute. Mellor's definitions of the softening range and softening point of a clay are not, however, commonly adopted, and ceramists almost invariably regard t,° C. as the softening point. Although the actual softening temperature may be measured by means of a pyrometer, it is more customary to express the result by giving the number of the Seyer cone that has the same softening point when heated alongside the clay under examination. Seger cones were originally composed of suitable mixtures of washed Zettlitz kaolin, Roistiand felspar, Norwegian quutz, Carrara marble, and pure ferric oxide; they were numbered from 1 to 39 in the order of increasing softening points. Cramer and Hecht, by adding boric acid and lead oxide to the list of ingredients, added the more fusible cones, numbered 01, 02, 03 . to 022; the softening points associated with the Seger cones then ranged from 600° C (cone 022) to 1880° C (conc 39) 2

The bending points of Seger cones only correspond to definite temperatures when the conditions of heating are those for which the cones were designed. Thus, while cone 10 can be observed to soften at about 1300° C when heated in a small gas muffle, yet when its temperature is very slowly raised in a large kiln, taking about four days to reach its maximum temperature, the same cone softens below 1200° C. Agam, a fireday that softened at cone 33 when heated in a standard manner, only softened at cone 35 when heated in a rapid injector furnace. For this reason Seger was always strongly opposed to the use of a temperature scale in connection with his

to lose their angularity

2 In 1908 it was decided to suppress cones 21 to 25, since their softening points were
very close together, and to modify the compositions of a number of the other cones. The
numbers by which the new Seger cones are designated, and the softening temperatures (in
degrees C) usually associated with them, are as follows (see Spiechsaul, 1908 p. 561).—

| 022   | 600° | 012a  | 855  | 02 α | 1060 | 9  | 1280 | 19 | 1520 | 34 | 1750 |
|-------|------|-------|------|------|------|----|------|----|------|----|------|
| 021   | 650  | 011 a | 880  | 01 a | 1080 | 10 | 1300 | 20 | 1530 | 35 | 1770 |
| 020   | 670  | 010 a | 900  | 1 a  | 1100 | 11 | 1320 | 26 | 1580 | 36 | 179  |
| 019   | 690  | 09 a  | 920  | 2 a  | 1120 | 12 | 1350 | 27 | 1610 | 37 | 182  |
| 018   | 710  | 08 a  | 940  | 3 α  | 1140 | 13 | 1380 | 28 | 1630 | 38 | 185  |
| 017   | 730  | 07 a  | 960  | 4 a  | 1160 | 14 | 1410 | 29 | 1650 | 39 | 188  |
| 016   | 750  | 06 a  | 980  | 5 a  | 1180 | 15 | 1430 | 30 | 1670 | 40 | 192  |
| 015 a | 790  | 05 α  | 1000 | 6 a  | 1200 | 16 | 1460 | 31 | 1690 | 41 | 196  |
| 014 a | 815  | 01a   | 1020 | 7    | 1230 | 17 | 1480 | 32 | 1710 | 42 | 200  |
| 013 a | 835  | 03 a  | 1040 | 8    | 1250 | 18 | 1500 | 33 | 1730 |    | 1    |

<sup>&</sup>lt;sup>3</sup> Mellor, Collected Papers, vol. 1. p 379. See also Zimmer, Trans. Amer. Cer. Soc., 1899, r, 23.

<sup>&</sup>lt;sup>1</sup> See Mellor, Collected Papers, vol 1 p 190. The Institution of Gas Engineers defines the softening point as the temperature at which the angular edges of the test pyramid begin to be a three angular edges.

cones. In general, the softening temperature of a clay is raised by increasing the rate of heating and it is decreased by increasing the pressure.

It has been previously mentioned that the colour of a raw clay is no safe criterion of the coloni that the same clay develops when heated. In , general it may be said that clays free from non burn white, those containing 2 or 3 per cent of non-oxide burn to a buff coloured mass, and those containing 4 to 5 per cent or more of non oxide yield a red product. The colour of a burned clay does not, however, depend merely upon the quantity of iron present, but is influenced by the size of the particles and the state of oxidation of the non, the presence of other substances, and the temperature to which the clay has been heated. When all the iron present in the burned clay is in the ferric state, the tendency is for the colour to pass from white through buff to red as the amount of non mereases, the colour deepening as the temperature of firing rises. Calcareous clays only burn to a buff colour even when much non is present, the bleaching action of the lime being greatest when its percentage is three times that of the ferric oxide; for this reason chalk is often added to brick-clays so as to produce white or buff-coloured bricks. The presence of titame oxide is said to intensity the colouring power of the non. When the nature of the clay and the firing conditions are such that ferrous oxide is produced in the mass, vitification commences at a comparatively low temperature, with the production of a dark blursh-grey or black fusible alummosilicate containing from in the ferrous state

In the burning of ferriginous clays it is usually desirable to convert all the non into the ferric state before vitrification commences, in order that the clay shall burn to the desired colour and not vitrify too readily. To accomplish this it is necessary that the burning shall take place in an oxidising atmosphere; any carbonaccous matter must be completely burned out at a temperature not exceeding about 900° C, any ferric oxide which has been reduced by the carbonaccous matter must be re-oxidised, and any pyrites or siderite initially present must be reasted to ferric oxide. The complete oxidation of pyrites or siderite in a clay is a very difficult task, and clays containing pyrites seldom burn red, instead, a buff-coloured product is obtained with black spots scattered irregularly over its sinface and throughout its mass. Printes and siderite, however, are rarely found in sinface clays, owing to the fact that by weathering they are converted into limonate, they are frequently found in shales and fue clays, particularly those associated with coal-sening

When the complete oxidation of the iron has been accomplished the colour of the surface of the birned clay may be altered, if desired, by changing the atmosphere in the kiln from an oxidising to a reducing one. The well-known vitrified Staffordshire "blue" bricks are produced in this manner

The vitrification of clay is usually associated with a marked change in the volume of the mass; almost invariably a shrinkage occurs. The volume change actually observed is the net result of a number of such changes, which are due to different circumstances. As soon as the fluxes commence to melt, the hand produced draws the unfused particles closer and closer together. The diminution in volume which results from this depends upon the extent to which vitrification is allowed to proceed, and upon the porosity

<sup>&</sup>lt;sup>1</sup> See Mellor, Trans Eng. Cer. Soc., 1907, 6, 142, 1908, 7, 112; Collected Papers, vol. i. p. 122, on pyritiferous clays.

of the clay just previous to the commencement of vitrification; the latter depends upon the nature of the clay and the shrinkage it has already undergone in the drying period The volume shrinkage due to the elimination of porces is partially compensated for by certain expansions in volume, Practically all crystalline solids expand on fusion, and the fusion of such fluxes as mice and felspar is accordingly associated with an increase in volume amounting with felspar to about 6 per cent. Polymorphic change may occur in clays rich in free silica and may be associated with an increase in volume. This arises from the fact that silica exists in quite a number of polymorphic forms, of which those stable above 800° C (tridymite, christabolite, etc ) have specific volumes some 14 per cent greater than those stable at lower temperatures Thus, highly siliceous fire-bricks expand when fired The processes of solution which are operative throughout the period of vitrification also influence the volume of the mass, though only to a very slight extent

Clays vary very much with respect to the amounts of fire-shankage which they undergo Excessive shunkage leads to warping and cracking in goods manufactured from them, and should be corrected by the addition

of suitable refractory materials (see p. 121)

Kinds of Clays.—Clays may be divided into two groups, refractory clays and non-refractory clays, according to whether their softening temperatures are above or below a certain arbitrary point, for which Mellor 2 proposes 1500° C., while the German ceramic chemists 3 and the Institution of Gas Engineers adopt the softening point of Seger cone 26 (about 1580° C )

Certain refractory clays become white or cream-coloured when fired. Of these, the chimi-clays and hadins are the most important, the other clays of this type are called pipe clays. The other refractory clays, which become grey, bulk or red in colour, are known as friedays. Mellor 2 divides them into three grades (1) low grade, softening between 1500° and 1650°, (11) medium grade, softening between 1650' and 1750'; and (iii) high grade, softening above 1750°. The Institution of Gas Engineers recognises two grades (1) No. 1, when no softening occurs below Seger cone 30 (1670°), and (ii ) No 2, when softening occurs above cone 26 (1580°), but below cone 30 (1670°)

The non-refractory clays may be either (1.) calcarcous or (11) non-calcareous, according to whether they do or do not contain more than about 5 per cent. of admixed calcium carbonate. The calcareous clays are usually termed malms (or marls), and, as already explained, when burned in an oxidising atmosphere they yield cream, buff, or red products according to the amounts of ferric oxide and lime present. The non-calcareous clays vary greatly in their properties, from the valuable ball clays which burn to a white or cream colour, to the more common buff and red-

<sup>1</sup> See Vol. V.

<sup>&</sup>lt;sup>2</sup> Mellor, Collected Papers, vol. i. p. 190.

<sup>3</sup> Cramer, Tonind. Zeit., 1902, 26, 1064.

<sup>4</sup> Many pipe-clays are not refractory, this term being extensively employed for any white clay which can be used for whitening belt;, etc. If a pipe-clay is found to be refractory it is at once classified among the china-clays or ball-clays, as these are of greater

The term "pipe-clay" is also applied to clays suitable for the manufacture of drain-pipes. Such clays are better considered as fireclays or ball-clays according to their nature; they are distinguished from the true pipe-clays in not being white when dry.

burning clays used in the manufacture of bricks, terra-cottas, and common pottery water

Calcarcous class in which the calcium carbonate is present in a finely divided state, i.e. as chalk, are largely used for making light-coloured bricks. Clays may be successfully used even though they contain 25 per cent, of chalk, provided that the product is only slightly vitrified. Calcarcous clays are also used in the manufacture of Portland cement.

China-lays and kiolius are used in the manufacture of china, porcelain, stoneware, fine faience, and ultramatine (see p. 135), they are also used for weighting paper 2. Some account of their nature has already been given (p. 104), and for further information the reader must refer to special treatises. Important deposits of china-clay are found in Cornwall and Devon in England; knolins occur at St Yriers, near Limoges, in France, at Zettlitz in Bohenna, in various parts of the United States, etc.

Ball clays are highly plastic transported clays which burn to a white or cream colour. They frequently contain 3 or 1 per cent, or even a larger proportion, of organic matter, and hence in the raw state they are often nearly black in colour. In England, the best deposits are found in Devenshine and Doiset. The ball clays form the basis of most ordinary pottery. When carefully selected they are extremely fine in texture and free from guit, and do not differ very much in composition from commercial clause. Proceedings are inferior qualities of ball-clay, containing rather more non and alkalies and considerably more shear.

Fireclays -The chief freelays are the clays and shales found in association with coal seams. In the British Isles such clays are found in large quantities in Northumberland, Durham, Yorkshire, Northughamshire, Derbyshire, Staffordshire (near Stourbridge), Warwickshire, Shropshire, North and South Wales, South-West Scotland (near Glasgow), and County Tyrone in Ireland The fireclays vary very much in composition, but all are relatively low in fluxes and to this fact they owe their refractorness. The shales occur above the coal seams, the non-lammated fireclays below them (underclass) Fireclass are generally grey or greenish grey in colour, owing to the presence of vegetable matter and non compounds (notably pyrites, which is detrimental to the quality of goods made from the clays), they are compact, dense, and of varying degrees of hardness By exposing them to the weather most firelays crumble and become easier to grand They are utilised in the manufacture of fire-bricks and other ceramic products of a refractory nature.

Brick-clays.—Although at times it has been convenient and economical to manufacture ordinary building bricks from frieclays, yet the greater part of the bricks and tiles used for constructional purposes are made from common, non-refractory, plastic clays which develop buff or red thits when fired. Common bricks may be satisfactorily made from clays so poor in "true clay" as to be scarcely worthy of being ranked as clays. Such clays, which are frequently termed brick earths, are usually rich in sand. So long as the adventitions materials present in a clay are chiefly sand and chalk, and provided that the clay is reasonably plastic and will commence to vitrify at about 950° C, with a volume contraction not exceeding 25 per cent, to

30.4

<sup>3</sup> See note 4, p. 118,

<sup>1</sup> See Vol III.

<sup>&</sup>lt;sup>2</sup> See Remington, Bowack, and Davidson, J. Ind. Eng Chem., 1911, 3, 550.

form a mass having a crushing strength of at least 85 tons per square foot, it will generally be suitable for the manifacture of common bricks

The clays utilised in the preparation of roofing-tiles, architectural and decorative terra-cotta, and coarse pottery wave are closely allied to those used for buckmaking, but are decidedly finer in texture, particularly those used for making the terra-cottas. When suitable clays are not found locally, it is sometimes convenient to prepare them from brick-clays by removing the coarsei particles by washing and sedimentation. Some fine-grained shales are also excellent materials for use in the manufacture of terra-cotta.

Fuller's earth is an earthy deposit similar in character to clay. It usually has a characteristic greenish-grey tint. Apart from its ready fusibility, it behaves like a china clay of unusually low plasticity. Its mineralogical composition is not yet clucidated, and the recognition of true fuller's earth is not at all an easy matter. In England the best deposits occur in the neighbourhood of Reigate (Surrey) and at Woburn Sands in Bedfordshire. The following data relate to the analysis of a sample from near Reigate —  $^2$ 

Fuller's earth absorbs grease, and was originally employed for fulling or degreasing wool. Its use for this purpose has declined considerably, but it is also used for clarifying and deodorising fats, oils, and greases (particularly laids, castors, coconuts, and cottonseed oils), as a diluent for ultramarine, and in the manufacture of paper, toilet powder, and certain soaps.

Ceramics. Classification of Ceramic Products. 1—For the production of ceramic ware a material is required that shall be (1) sufficiently plastic when moist to be formed into any desired shape, and (n) dense and stone-like when "fired," i.e. heated to a sufficiently high temperature. Accordingly, the foundation of every piece of ceramic ware is one or other of the various natural clays, or a mixture of them. For many purposes, however, it is impossible to use clay alone, and other suitable materials have to be added to it. The clay may, for instance, be so plastic that it is unable to retain its form during the process of drying, or it may become deformed or develop cracks owing to excessive shrinkage during the same process. To overcome these and other defects it is necessary to add hardening or opening materials to the clays, and thereby modify the plasticity and shrinkage of the mass. Again, in certain kinds of pottery it is essential that the fired body be impermeable to water, and it may then become necessary to add considerable quantities of suitable fusible materials or flines to the clay.

It appears to be composed mainly of one or two of the minerals related to clay.
 Analysis communicated by Mi A. G. Hopkins

<sup>&</sup>lt;sup>2</sup> Analysis communicated by Mi A. G. Hopkins
<sup>3</sup> The reader seeking further information on the ceramic industries is referred to the following works. Boutry, A Treatise on Ceramic Industries, revised translation by A. B. Searle (Scott, Greenwood & Son, 1911), Grangert La ceramique industrielle (Gauther-Villars, Paris, 1905), Searle, Modern Brickmaking (Scott, Greenwood & Son, 1911), Granger, Fabrication et emploi des matériaux et produits réfractaires utilités dans l'industrie (Paris, 1910); and the works of Thorpe, Moissan, and Le Chatcher eited on p. 103. A fairly complete list of books on ceramics is given in Searle, Clayworker's Handbook (C. Griffin & Co., Ltd., 1911).

Accordingly, so far as the preparation of pottery bodies is concerned, the materials used may be divided into three groups, as follows

(i.) Plastic Materials—The different varieties of clay

(ii) Opening or Hardoning Materials - These include silica in the form of sand, finit or quarts, "grog" (i.e. previously fixed clay), and a number of other materials less frequently used

(ni) Fusible Materials of Fluxes — Felspar, pegmatite, bone ash (i.e. calcium phosphato), and chalk are the fluxes most commonly employed. The felspar employed is usually Norwegian orthoclase. In England the pegmatito used is known as Cornish stone or china stone, it is a partially knolmised granite which behaves in many respects like a mixture of felspar and quartz. The bone ash is generally prepared by calciumg the bones of oven

The numerous products of the ceramic industry may be divided into two classes, according as the "bodies" are (i) permeable or (ii) impermeable to water. The bodies in class (i) are porous, exhibit an earthy fracture, and may be seratched by steel, those in class (i) are hard and sonorous, exhibit a viticous or conchoidal fracture, and cannot be scratched by steel. The bodies of class (ii) may be divided into two subclasses, (iii) procedures, which are white and translucent in pieces 2 or 3 mm, thick, and (ii) b) sto awares, which are opaque and usually coloured, if only to a slight extent

Porous ware may either be used as such for constructional or decorative purposes, e.g. numerous kinds of bricks and tiles, and terra cotta statuary, or it may be covered with a vitrous envelope known as a glaze or an enamel according as it is transparent or opaque. Porous pottery when glazed or enamelled is known as the carthenware when the body is white and as jained when the body is coloured, although the enamelled faience is sometimes called majolina. Unglazed porous ware may be divided into two subclasses, (i. a) those which lose then form or become imperimeable at a temperature higher than that at which hard porcelam is fired (c. 1400°C) and (i. b) those which do not. The former may be termed terra cotta, using the word in its literal sense (baked earth), and the latter, nicelay or retractory ware. Saintary ware is chiefly made by glazing articles made of fireday and is, therefore, to be included under frieday ware. When it has a vitified body it should be included under stoneware.

Impermeable ware, like porous ware, may or may not be glazed. Stone-ware is often glazed or enamelled, and most porcelains are glazed, although with the variety known as param this is seldon the case.

The preceding classification of ceramic products is that proposed by Bourry and is summarised in the following table —

- I. POROUS BODY, PERMEABLE TO WATER
  - (a) Not glazed terra-cotta
  - (b) Not glazed fireclay goods
  - (c) Glazed fine earthenware, sanitary ware, faience, majolica.
- II BODY IMPERMEABLE TO WATER.
  - (a) Translucent porcelain.
  - (b) Opaque stoneware

Ceramic wate may be glazed (or enamelled) either (1) by drying the "body" until it is in a fit state to have the glaze applied to its surface,

<sup>&</sup>lt;sup>1</sup> This distinction between the uses of the terms "faience" and "majolica" was proposed by Burt (Trans. Amer. Cor. Soc., 1904, 6, 109).

and then firing the mass, or (ii.) by firing the unglazed body, coating the "biscuit" ware thus obtained with the glaze, and re-firing. The two firings are said to be carried out in the "biscuit" oven and the "glost" oven respectively, and the second firing always occurs at a lower temperature than the first. The second procedure is nearly always adopted, but hard porcelain, some saintary ware, and certain kinds of stoneware are prepared by the first method.

The porcelams, and particularly the hard porcelams, are of particular interest and importance to the chemist. Being covered with hard glazes that are scarcely attacked by prolonged boiling with inneral acids or aqueous solutions of alkalies, and being capable of withstanding rapid alterations of temperature, they are invaluable for the manufacture of circubles, basins, etc. To the chemical manufacturer the coarser stoneware is of equal importance for the construction of chemical plant, e.g. condensors of mineral acids,

evaporating pans, storage tanks, pumps, taps, etc.

1. Terra-corras—In accordance with the classification given above, this term includes such articles as ordinary, hollow, ornamental, vitified, and light bricks, tooling and paving tiles, blocks, dram-pipes, and common unglazed pottery ware (such as flower-pots), in addition to the architectural and decorative wares to which the term terra-cotta is usually restricted. Such goods are prepared from natural clays, or suitable mixtures of clays, with the occasional addition of chalk or said (see p. 119)—Light bricks (or terra-cotta lumber) are obtained by incorporating sawdust or chopped straw with the clay used, the vegetable matter being, of course, burnt away in the firing process—In order to reduce the tendency to crack and to facilitate drying and burning, the larger pieces of terra-cotta are usually made hollow, and in the production of very large pieces a coarse porous clay is generally used to form the bulk of the mass, a finer terra-cotta clay being then used to cover the face of it

n. Fireclay Goods.—These are distinguished from all other ceramic products by their resistance to heat, they include bricks and blocks used for hining furnaces, muffle kilns, crucibles, retorts, etc, employed in the metallurgical, glass and ceramic industries. They are prepared from fireclays, with the addition, when increasily, of a suitable opening material such as grog (or chainotte) and silica, but as the latter reduces the refractoriness of a good fireclay it is preferable to use the former, which consists of fireclay that has been fired at a high temperature and ground to a coarse powder. The texture of the fired article may also be more readily readjusted by the use of grog than silica. On the Continent it is customary to use a large proportion of grog, often as much as two parts of it to one of raw clay, in forming the fireclay body, the maximum amount of grog permissible depends on the plasticity of the clay. In England the use of grog in the manufacture, of fireclay goods is not so extensive

The practical value of a fireclay article is not determined solely by its refractoriness, i.e. its ability to be intensely heated without softening. It may have to withstand sudden temperature changes, the action of flames or molten corrosive slags, etc., according to the manner in which it is used, and these points have to be considered carefully when deciding on the composition and the texture of the body. Thus, suppose firebricks are required to with stand abrupt temperature changes. It is found that the more porous the texture the less will be the tendency of the bricks to crack, and that without the addition of opening material raw fireclays do not fire to sufficiently porous

bricks. Accordingly, the fireclay used is mixed with the requisite amount of grog of a suitable degree of fineness, the coarser the grog and the greater the amount used the less the tendency of the brick to etack with sudden changes in temperature. For most purposes, however, firebricks should have a close compact texture so as to possess strength and resistance to abrasion.

Fireclay goods rich in alumina (e.g., bauxite bricks) are piepaied by opening up fireclays with bauxite (p. 73), which, to be satisfactory, must not contain more than 10 to 15 per cent of feiric oxide

Fireday goods extremely tich in silica, eg silica bricks and acid linings for steel furnaces, are prepared from gainster tock. Dinas tock, or crished silica. Gainster is a fine-grained siliceous grit, the composition of which approximates to time parts of silica mixed with one of fireday, for use as a refractory material it must be very carefully selected. Dinas rock occurs in Wales in the Vale of Neath, it contains 97 to 98 per cent, of silica and is practically free from clay. For making bricks 1 to 2 per cent of lime is added to act as a flux and the wet mixture is pressed into shape. Crushed silica is used on the Continent for making silica bricks, as no natural material of the quality of Dinas rock is available.

Carbon is used as an opening and heat conducting material in the preparation of plumbago crucibles. Either graphite or the deposits of carbon found in gas-retorts may be utilised, the proportion used varies from 25 to 50 per cent of the body mixture. Part of the fireday may be in the form of fine grog. Sometimes magnesia is added to the body.

in Earthenware and Fairness —For common pottery ware, architectural fairness such as glazed bricks, tiles, etc., engainste tiles, vases, dishes, and other decorative objects, the body consists simply of a plastic clay with the addition of sand or grog if necessary. The so called enamelled fairness, e.g., stove tiles and imitation Moorish, Italian, and French fairness, are prepared from clays naturally or artificially calcarrons, the body usually containing from 15 to 20 per cent of lime when fired. The domestic white or cream-coloured carthenware is prepared from a more complex body, the basis of which is ball-clay. To whiten the body china clay is added and to open it up flut is used, as a flux felspar or Cornish stone is utilised. To eliminate the pale yellow or cream colour from the fired body a little oxide of cobalt is usually added (0.02 to 0.08 per cent.). The following table shows how the relative proportions of the migredients of an earthenware body may be varied:—

| Ball-clay     | 50 | 47 | 13 | 31 | 25 | 25  | 24      | 21 | 18 1 | zer cent. |
|---------------|----|----|----|----|----|-----|---------|----|------|-----------|
| Chma-clay     | 10 | 24 | 21 | 36 | 32 | 25  | $^{27}$ | 28 | 43   | ,,        |
| Flint         | 35 | 22 | 23 | 21 | 34 | 4() | 36      | 38 | 24   | ,,        |
| Cornish stone | 5  | 7  | 10 | 12 | 8  | 10  | 13      | 13 | 15   | .,        |

French siliceous faiences, which form admirable artistic wares, are prepared from a body containing china-clay 21, chalk 24, fint 48, and frit 4 parts, the frit being piepared from Fontainebleau sand 85, potassium carbonate 7, sodium carbonate 3, and chalk 5 parts. This body was introduced by the celebrated faience manufacturer Deck in endcavouring to reproduce Persian faiences.

iv Stoneware — Fine English stoneware is prepared from a body similar in composition to that of earthenware, but the proportion of Cornish stone.

is increased and that of the flint decreased, as shown by the following

| Ball-clay .   | 10 | 25   | 25 | 29 | 45 | $29~\mu$ | er cent. |
|---------------|----|------|----|----|----|----------|----------|
| Chma-clay     | 15 | 12.5 | 20 | 14 | 15 | •        | 11       |
| Flut.         | 20 | 12.5 | 10 |    |    |          | 1)       |
| Cornish stone | 55 | 50   | 45 | 57 | 10 | 71       | 11       |

"Common" stoneware articles, e.g. paving tiles, drain-pipes, samtary ware, and chemical stoneware, are prepared from stoneware or vitrifiable clays (cf. p. 115) These clays when burned form opaque imperincable waie, usually of a greyish, yellowish, or blursh shade. They must be selected with great care, as stoneware clays are very hable to twist and warp when heated. The number of sources of good stoneware clay is not very considerable, the best of those found in England being the upper and less pure portions of the Devonshire and Doiset ball-clays and the more fusible fireclays Accordingly, stoneware manufacturers are frequently obliged to adapt other clays to their needs - For this purpose felspar or Cornish stone may be added to fireclay, or a refractory material may be added to a fusible non-calcareous clay containing clay or felspar v. Porcelain --The porcelains may be divided into two groups, (a) the

hard porcelains and (b) the soft or tender porcelains The hard porcelains are distinguished from the soft ones by their greater resistance to heat. When glazed, they are usually made so that the very hard glaze is fired with the body in one operation

The basis of all porcelain is china-clay or kaolin. In order to obtain a more plastic body a little ball-clay may be added. The increased plasticity, however, is only gained at the expense of the translucency of the finished wate, and ball clay is therefore usually excluded in the preparation of hard porcelain table-ware and ornamental goods, but utilised in the manufacture of porcelain for making electrical insulators or liming ball-mills and other vessels

In the preparation of hard porcelain the opening materials used are quartz and biscuit porcelain, and the fluxes employed are felspar, chalk, and gypsum From the point of view of the chemical composition of the products, the hard porcelams may be divided into those rich in alumina and very suitable for chemical purposes, and those rich in silica and well adapted for the preparation of ornamental ware. The following bodies are employed in the preparation of a number of hard porcelains -

|        |   |    | Kaolin   | ' E       | Pelspar         | . !                 | Quartz.                   | Chalk               | Gypsum                     | Biscuit<br>Porcelan        |
|--------|---|----|----------|-----------|-----------------|---------------------|---------------------------|---------------------|----------------------------|----------------------------|
| rcelam |   |    | 65       | -         | 15              | -                   | 14                        | 6                   |                            |                            |
| 1)     | : |    | 75       |           | 25              |                     | ••                        | Ĭ                   | ':                         | _                          |
| ,,     |   |    | 73<br>72 |           | $\frac{25}{12}$ |                     | 12                        |                     |                            | 2                          |
|        | • | ., | .,       | rcelan 65 | rcelan 65       | roelain . 65 , 15 , | rcelain 65 15 75 25 73 25 | rcelam . 65 , 15 14 | rcelain . 65   15   14   6 | rcelain . 65   15   14   6 |

Presumably, the kaolins used include certain amounts of quartz and felspar or mica (see p. 105) The three most important constituents of hard porcelain are kaolin, quartz, and felspar, and according to Vogt 1 their relative propor

<sup>1</sup> See article "Céramique" in Moissan, Traité de chimie minérale (Paris, vol. iv., 1905).

tions should be kept within the following extremes kaolin 65 to 35, quartz, 15 to 25; and felspar, 20 to 40 per cent.

The only kind of soft porcelain of any industrial importance is that which is variously known as phosphatic potcelain, English potcelain, hone china, or simply china, this china nare is almost evelusively produced in England. Its special characteristic is the large proportion of bone ash (30 to 50 per cent.) which enters into the composition of the body, which is made chiefly from china-clay, Cornish stone, and bone ash. The limits of composition are very wide, the following may be cited as examples of suitable mixtures.—

| Chma-clay .  | 25    | 30 | 22 | 29 | 34 p | er cent |
|--------------|-------|----|----|----|------|---------|
| Cormsh stone | 25    | 35 | 31 | 30 | :::  | ,,      |
| Felspar      | • • • |    |    |    | 19   | "       |
| Bone ash     | 50    | 35 | 47 | 11 | 47   | ••      |

Parian is a soft porcelain which is usually unglazed. The body consists essentially of china-clay and felspar, a very large proportion of the latter being present, and in the preparation of English parian these appear to be the only substances used, the body containing 33 to 37 per cent. of chinacilay. Parian wave closely resembles fine marble.

Information concerning the other varieties of soft porcelain, e.g. French or fritted porcelain, new Sevres porcelain, and Seger's porcelain, must be sought in special treatises on ceramics (see footnote on p. 120)

Preparation of Ceramic Bodies.—The various ingredients that go to form the body of a ceramic product must be reduced to a state of fine division, and thoroughly and uniformly mixed. In the cases of cheap articles, e.g. bricks, where a natural clay alone is required, the preparation of the bodies is comparatively simple, it being frequently sufficient to crush or remove stones and other hard foreign bodies and then pass the wet clay through a pug-mill to secure the necessary uniformity. Some clays are dried and crushed to fine powder, all shales must be crushed and ground. The greatest care in the preparation of uniformly plastic bodies is naturally required where several different ingredients have to be ground and mixed, eg in the preparation of fine earthenware, stoneware, and porcelam In English fine earthenware, for instance, the china-clay, ball-clay, china stone, and flint having all been reduced to the state of fine powder suspended in water, the requisite quantities of the "slips" are run into a large vat and thoroughly agitated. The "slip" thus obtained is passed through fine sieves to remove coarse particles and run over permanent or electro-magnets in order to remove particles of iron and magnetisable iron compounds. It then passes to filter presses, where most of the water is removed by filtration through canvas, and, the plastic body being harder on the outside than on the inside, it is thoroughly beaten up or put through a pug-mill to render it uniform and free it from airbubbles Instead of adopting the preceding method of "shp-blending," the raw materials are sometimes ground and dried; the requisite quantities are then weighed out and mixed with water.

Shaping Ceramic Articles.—Of the various methods by which the plastic body is formed into shape, the oldest and most general is that known as "throwing on the wheel" R is used in the preparation of round articles from sufficiently plastic bodies. "Turning" on a lathe is also adopted, particularly when an article must be made very accurate in shape and size. A process known as "expression" is used in the production of drain or sewer

pipes and building blocks, bricks and tiles. For this purpose, the plastic paste is forced through a die or template so that the issuing clay is of the desired section; it is then cut into suitable lengths. The better qualities of bricks and interlocking roofing tiles are largely made by pressing the stiff plastic paste in metal moulds by powerful presses, and the same process is sometimes applied to bricks and tiles which have been roughly moulded and allowed to become half dry "Dry pressing" is used in the production of certain classes of ceramic ware, particularly tiles used for internal decoration, the body is dried, crushed to powder, damped until it adheres together loosely when squeezed, and filled into a metal mould, where a die is forced down upon it by hand or mechanical power.

The simplest method of moulding a plastic body is that used in making bricks by hand with the use of wooden moulds. More ornamental articles are formed by beating the plastic body into a thin cake or "bat" and pressing it into moulds made of plaster of Paris. Frequently a mould has to be made in sections, which are fitted together and strapped up, the joints being smoothed down so that the clay forms one piece inside the mould. The clay dries a little on standing, contracts slightly, and detaches itself from the mould, the portions of which are then removed. "Shp casting" is another method of moulding, particularly suited to feebly plastic bodies such as porcelains. The "slip," prepared of such a consistency that one pint weighs 30 to 36 ounces, is poured into the plaster mould. The plaster absorbs the water from the slip in contact with it and a coating of the body is formed on the inside of the mould. When the coating is sufficiently thick the remaining slip is poured out and the mould and its contents dired in a hot stove. By this process it is possible to form teapots, jugs, etc., in one piece, instead of having to form them in parts and join these together afterwards.

The ware having been formed, it is slowly dried in an open shed,1 in a drying room, warmed by waste heat from the firing kilns, or by other suitable means. With some porcelain bodies, however, the final temperature reached in drying is much higher (700° to 900°)

The ware is then ready for firing, unless it is to be fired and glazed at one operation, when it must first receive its coating of glaze

Firing.—Ceramic bodies are usually fired in kilns or ovens.2 Such products as bricks, tiles, drain-pipes, etc, may be exposed directly to the flames, but this is not as a rule permissible with wares of better quality, which must therefore be packed in fireclay boxes, known as saggars. The ovens are large brickwork chambers which may be likened to vertical reverberatory furnaces. The saggars are stacked in the ovens in piles or "bungs" and the flames allowed to play directly on to them

The final temperature reached in the kiln and the duration of the heating depend largely upon the nature of the ware required. Ordinary bricks, for instance, may only require heating to about 950° C. until vitrification, produced mainly, as a rule, by the fluxing action of lime, has so far advanced that the fired product has the necessary crushing strength. Hard porcelain,

<sup>1</sup> During the summer months bricks and coarse ware may be dried in the open air, if

protected from the sun and wind.

There are many types of kiln and oven, each suited to the production of a particular kind of ware. A description of these is out of place in the present volume, and the reader requiring further information is referred to the treatises mentioned in the footnote on p. 120.

on the other hand, must be fired at 1400° C. or thereabouts and the vitrification pushed almost to the extreme limit

It has already been mentioned (p. 121) that fired ceramic bodies may be divided into two classes, the permeable and the impermeable. In practice, however, these classes cannot be divided rigidly. The fine English white wares, in particular, are difficult to classity, the distinction between some pieces of faience (carthenware) and some of stoneware being almost impossible to define. Thus, the term "stoneware" is frequently applied to glazed pipes, tiles, and saintary ware, the bodies of which are decidedly porons though the articles as a whole are impermeable

In discussing the changes that take place in ceramic bodies during the firing process, it will only be necessary to refer to friedly goods, white ware and porcelam, the nature of the changes undergone by other bodies having been sufficiently indicated in dealing with the action of heat on clays (p. 113).

In the preparation of firebricks it is of great importance that the goods shall be fired long enough and at a temperature high enough to prevent any marked after-contraction when the bricks are in use. Otherwise, under favourable temperature conditions, contraction will continue at an appreciable rate and give rise to movements in the masonry of the furnace and setting, The choice of the fireday should be influenced by the maximum temperature likely to occur in the finance in which the bricks are to be used. Allowing a reasonable margin of safety, the most suitable softening temperature for the clay may be obtained, and an unnecessarily refractory clay should be rejected as being likely to produce bricks too open in texture and correspondingly weak mechanically Non-vitrifiable fireclays, poor in fluxes, exhibit a maximum contraction on firing, this varies with the temperature and rate of heating. When the maximum contraction has been attained, subsequent herting at the same temperature is attended by a slight expansion and a slight disintegration of the brick 1, subsequent heating at a higher temperature, however, produces a further contraction, which attams a maximum as before

As fireclay goods are manufactured for the purpose of withstanding high temperatures for prolonged periods, a few words on the nature of the changes they undergo on continued heating may be added. The ultimate character assumed by a fireclay body is that of a nearly homogeneous vitreous mass, and as this condition is approached the body tends more and more to soften and lose its shape. The longer it takes to realise this state of affairs, the longer is the life of the fireday body. With prolonged heating there is usually associated a decrease in the crushing strength of the body. This appears to be due to a number of causes, one of which is the gradual loss of alkalies by volatilisation, followed by a slight disintegration of the "cementing' matrix which binds together the solid particles of the body 2 Another cause is the separation of crystals of sillmanite from the vitreous matrix. The rate at which this crystallisation proceeds doubtless depends upon the proximity of the temperature to the best crystallising temperature of the matrix, as explained later in connection with glazes (p. 131) The crystallisation being irregular, and attended

Mellor, Collected Papers, 1914, vol 1. pp. 309-315.
 Cramer, Tomand Zett., 1887, 21, 288, Mellor and Austin, Trans. Eng. Cer. Soc., 1907, 76, 129, J. Soc Chem. Ind., 1907, 26, 375, 380
 Microscopic examination reveals the growth of local patches of crystals.

by a contraction in volume, local strains are set up in the body and tend to rupture it.1

In the firing of hard porcelain 2 the felspar melts to a liquid which binds together the particles of heated clay and silica. As the heating is continued the solid particles are slowly attacked and dissolved by the molten flux, the clay more readily than the free silica, at about 1300° felspai can dissolve about 20 per cent of china-clay and 15 per cent of silica. The translucency of the fired body may be thus explained, the body consists of a transparent glass in which particles of silica and heated china-clay are embedded, forming a kind of framework in it Now, china-clay particles are permeated with minute capillary pores, and the particles left after the water of constitution has been expelled still retain this porous structure. When these pores are filled up by a colourless liquid, china-clay or the material left after heating it to redness is found to be fairly transparent. The translucency of porcelain is thus a result of the molten flux penetrating into the capillary pores of the heated clay at the same time as it is slowly dissolving the clay, and the translucency mercases in proportion as the percentage of china-clay in the porcelain body is reduced. It is clear, however, that there must be at least a certain (minimum) proportion of clay present, to ensure that the body shall not collapse when fired owing to lack of sufficient clay-silica framework.

Patches of accoular crystals of sillinamite frequently separate out from the glassy matrix when hard porcelain is fired, and, indeed, they seem to be produced whenever mixtures rich in clay and felspar are fired at or above 1300° C.3 The rate at which the crystals develop depends upon the temperature, and their production in heated porcelam probably explains the development of brittleness in porcelam tubes which have been heated for a considerable length of time.

The slight bluish tint usually noticed in porcelain is due to the body always containing traces of iron and titanium and to the final heating and vitrification being carried out in a slightly reducing atmosphere gives place to a vellow tint when an oxidising atmosphere is used

Owing to the large amount of flux present and the extent to which vitrification is allowed to proceed, porcelain contracts very considerably (about 15 per cent) in the fining. In soft parian porcelain, where the percentage of flux is extremely large and opening material is omitted, the contraction is still greater. Since parian is fired at a much lower temperature than hard porcelain 5 it does not show well-developed crystals of sillmanite unless, as is sometimes the case, a little lead glass is added to the parian The introduction of the lead glass appears to accelerate the rate at which the sillmanite crystals form

From what has already been said the nature of the changes produced when bone china is fired may be inferred, especially as powdered bone ash

For further information concerning the action of heat on friedlays see Mellon, J. Soc Chem. Ind., 1907, 26, 375, Mellon and F. J. Austin, low cit., Mellon, Collected Papers, vol. 1, pp. 309, 316, 379
 Hussak, Sprechaul, 1889, 22, 136, 154, Mellor, Trans. Eng. Cet. Soc., 1909, 5, 75, J. Soc Chem. Ind., 1907, 26, 375, Mellon and Austin, low cit.; Heath and Mellor, Trans. Eng. Cer. Soc., 1907, 7, 80; or Mellor, Collected Papers, pp. 19, 43, 98, 138
 W. Vermackk, Bull. Soc. franc. Min., 1889, 12, 447, 1890, 13, 256; Hussak, low cit.; Mellor, low cit.
 See later, under Glazes, p. 131.
 Haid porcelain is fired at 1250°-1400° C., painan at 1150°-1200° C.

has a porous structure similar to that of china clay, and so assists in producing the translucency of the china. Bone china is fired at 1250"-1300" (', and when correctly fired it approximates to a homogeneous vitrified mass more closely · than almost any other type of pottery. The temperature has to be regulated with extreme care, as when over heated, the china begins to lose its shape, phosphoric oxide fumes are evolved, the body acquires a sponge like character and "bloats," The loss of phosphorus is due to the presence of carbon or other reducing substance in the body or in the kilicat certain stages of the firing 1

The changes which take place when fine stoneware and white earthenware bodies are fired are similar to those which occur in the firing of hard porcelain, but vitrification is not allowed to proceed to the same extent. Less flux is therefore required and the firing temperature need not be so high. Fine stoneware has the same general structure as hard porcelam, but it is not translucent Why the substitution of ball clay for china clay should destroy the translucency of porcelam is not definitely understood, but it is probable that owing to the comparative ease with which the impure clay softens, the pores become closed up before the molten flux can penetrate into them

Glazes and Enamels.—Glazes and enamels are viticous envelopes used to cover ceramic bodies, either for decorative purposes or for rendering them impermeable to liquids. The finely divided glaze is spread over the surface of a ceramic body and the whole heated until the glaze melts and forms a thin film covering the surface continuously. The glazed body is then allowed to cool, when the molten glaze solidities to an amorphous "glass'

A pottery glaze should be clear, brilliant, and thin, so as not to hide modelled work, it should soften more readily than the body to which it is applied, and while being sufficiently viscous when molten not to run off an upright object it should be mobile enough to flow smoothly over the whole surface, it should be very resistant to attempts to devicinfy it, its coefficient of expansion should agree with that of the body in order that it may not chip off from edges or develop minute cracks (known as crazes) upon the surface, it should be unattacked by water and acid vapours, and for culmary, domestic, or sanitary purposes it should be resistant to dilute acids

Glazes vary very much in chemical composition and in firsibility all contain silica, alumma, and one or more of the oxides of the alkali or alkalino carth metals as necessary constituents. Of the other basic oxides that may be present, lead oxide is the commonest, while many of the more fusible glazes contain a second acidic oxide, namely boric oxide. In ceramics the compositions of glazes are usually expressed in a conventional manner by indicating the relative numbers of formula-weights of the various oxides present in such a manner that the total number of formula-weights of the oxides of um- and bi-valent metals is unity, the following, for instance, represents a typical earthenware glaze -

$$\left. \begin{array}{c} 0.25 ({\rm K\,Na})_2 {\rm O} \\ 0.35 ({\rm aO} \\ 0.40 {\rm PbO} \end{array} \right\} \\ 0.2 \ {\rm Al_2O_4} \left\{ \begin{array}{c} 3.0 \ {\rm SiO_2} \\ 0.4 \ {\rm B_2O_3} \end{array} \right.$$

The general glaze "formula" may be written RO JRO, yRO, when it is found that y is always greater and x is generally less than unity In general, the more bases of the RO group there are present, the more fusible the

<sup>1</sup> B. Moore, Trans Eng Cer Soc., 1905, 5, 37.
2 Or, more accurately, a mixture of substances which react to produce the glaze when heated.

glaze; replacement of silica by boric oxide also increases the fusibility. The proportion of alumina present largely influences the properties of a glaze, particularly its viscosity and its tendency to devitify. The use of lead in glazes has been known and practised for centuries. Lead glazes are very quick to mature and, by suitably choosing their compositions, may be adapted for use with a fairly wide range of pottery bodies. They are not suitable, however, for use with hard porcelain and once-fired stoneware, since lead oxide volatilises rapidly at the firing temperature and, moreover, the glazes mature too rapidly. The use of lead glazes is attended with certain tisks, owing to the poisonous nature of lead compounds. Lead-poisoning may affect the workpeople engaged in handling the glazes, or it may affect the purchaser of a lead-glazed culmary article, unless the latter will resist the action of boiling, concentrated acetic acid. The latter evil, however, has not ansen in England

Glazes containing only the constituents already mentioned are colourless and transparent. They may be converted into white, opaque glazes or enamels by the addition of stamme oxide or zine oxide in excess of that required to saturate the glaze. The opacity is due to the presence of solid oxide of zine or tim in suspension in the glaze. Calcium phosphate is sometimes used as an opacifying agent.

In general, the glaze is produced upon the body by the interaction of a number of suitable raw materials under the influence of heat. When all the raw materials are insoluble in water it is sufficient to grind them together and apply the mixture to the body, but when some of the raw materials are soluble (e.g., sodium carbonate, borax) a different procedure is adopted. The soluble ingredients are mixed with suitable proportions of the others and the mixture melted into a glass or firt. The first is cooled, powdered, and mixed with the remainder of the insoluble ingredients.

The glaze inviture may be applied to the body in various ways. It is commonly applied by "dipping," for which purpose the glaze mixture is stirred up into a "ship" with water and the article to be glazed then dipped into it. The body, owing to its porosity, absorbs a certain amount of water and a film of glaze mixture is deposited upon its surface. Other methods include pouring a suspension of the glaze mixture over the body as it is rotated, and spraying the suspension over it.

A glaze should be regarded as a supercooled, highly viscous, and more or less imperfectly inved solution. Accordingly, although a glaze may be made up in several different ways so as to have the same ultimate composition, it does not follow that the final glazes will "mature" in the same way. The behaviour of the glaze varies with the manner in which it is compounded, the physical condition of its constituents, and the character of the body on which it is fired.\(^1\) A glaze, being amorphous, has no real melting-point, but becomes less and less viscous as the temperature rises. Eventually, it begins to lose its shape and begins to flow like a mobile liquid. It is difficult to determine a precise temperature at which this occurs, but a small temperature range may be located fairly readily and the mean temperature spoken of as the softening temperature of the glaze. The softening temperatures of lead sheate glazes have been measured by Mellor, Latimer, and Holdcroft,\(^2\) who found that the gradual addition of more and more silica rapidly lowers the softening-point

Mellor, Trans. Eng. Cer. Soc., 1913, 11, 1, Collected Papers, vol. 1 p. 339.
 Mollor, Latimer, and Holdcroft, Trans. Eng. Cer. Soc., 1909, 9, 126, Mellor, Collected Papers, vol. 1. p. 247.

State of a

from 877°C (the melting-point of litharge) to 526°C. (for PbO+0·21SiO<sub>2</sub>) and then gradually increases it until with a composition corresponding to PbO+2·4SiO<sub>2</sub> the softening-point is 661°. All the softening points are much lower than the corresponding melting-points in the system litharge silica.

A pottery glaze, being a supercooled solution, has a certain tendency to crystallise. It has been found that the rate at which a supercooled hquid crystallises increases with the degree of supercooling up to a certain point, when the further augmenting of the degree of supercooling leads to a steady diminution of the rate of crystallisation. Hence in the cooling down of a glazed pottery body there is a certain temperature at which crystallisation of the glaze, when once started, proceeds at its maximum rate. Since in general it is not desired that the glaze shall crystallise, the temperature of the body should be hurried past this "best crystallising temperature," as all the time it is in the vienity of it the body is in a kind of danger-zone. The length of time that the body is in the danger zone depends upon the rate of cooling and also upon the rate at which the velocity of crystallisation diminishes as the temperature changes above or below the best crystallising temperature.

A pottery glaze cannot be looked upon merely as an envelope for the body, the union being much more intimate. All the time that a glaze is maturing it is attacking the surface of the body. Where the glaze and body meet there is thus formed a concentrated solution of body in glaze, and owing to the high viscosity of this solution and of the glaze they have not sufficient time to become uniformly mixed. Consequently crystals sometimes separate out from this intervening layer of solution on cooling, and cause the glaze to peel off from the body.

Although the crystallisation of a glaze is usually a defect, the production of crystalline glazes can, in certain oncumstances, be utilised for the purposes of pottery decoration. For instance, the intentional crystallisation of zinc silicate from glazes may be made to produce extremely beautiful effects "Sunstone" and "aventume" glazes are other well-known crystalline glazes, in which the crystals which separate out are complex silicates somewhat similar to the micas.

Hard porcelain glazes are rich in silica and alumina and comparatively poor in bases, the only bases present are potash, soda, lime, and magnesia. At Sevres the hard porcelain glaze consists of pegmatite from St Yilex, in Austria and Germany mixtures of felspar, quartz, and kaolin are used, with the addition, at times, of calcium carbonate or sulphate. The following are some of the compositions which have been published.

| Felspar     |       |  |      | 32.5 |      | p    | er cent |
|-------------|-------|--|------|------|------|------|---------|
| Kaolin      |       |  | 30 Q | 95   | 37.5 | 37.0 | ,,      |
| Quartz      |       |  | 440  | 34.5 | 50 0 | 37.0 | "       |
| Chalk       |       |  |      | 120  | 12.5 | 17.5 | "       |
| Gypsum      |       |  | 135  |      |      |      | "       |
| Biscuit por | celam |  | 125  | 11.5 |      | 8.5  | **      |

Chmese and Japanese porcelain glazes are also calcareous. The compositions of porcelain glazes fall within the limits RO  $0.2R_2O_3:3.5RO_2$  and RO  $1.25R_2O_3:12RO_2$ 

See Meller, Collected Papers, vol. 1 pp 6, 47, on Trans Eng Vrs. Soc., 1905, 4,
 J. Soc Chem. Ind., 1907, 26, 375, Ruddle, Trans Amer. Cer. Soc. 1906, 8, 326
 On crystalline glazes see W. Burton, J. R. Soc. Arts, 1901, 49, 213, 1904, 52, 595.

In the porcelains there is a very intimate relation between body and glaze, since they are fired together and are similar in composition. This is not so much the case with bone china glazes, which are applied to biscuit ware, usually contain both lead and boric acid, and are very similar to the glazes applied to fine stoneware and earthenware. The following are stated by Burton to be typical glaze recipes, (a) for china ware, (b) for white earthenware, and (c) for granite ware

|                                              | Glaze.                |                                |                              | 1                                                             | Fru                               |                       |                                    |
|----------------------------------------------|-----------------------|--------------------------------|------------------------------|---------------------------------------------------------------|-----------------------------------|-----------------------|------------------------------------|
| Frit<br>Flint<br>Cornish stone<br>White lead | (a)<br>45<br>15<br>15 | (b)<br>230<br>60<br>160<br>120 | (c)<br>360<br>50<br>50<br>80 | Borax .<br>Chalk<br>Fluit<br>Cornish stone<br>Felspai<br>Soda | (a)<br>50<br>10<br>40<br>24<br>33 | (b)<br>48<br>38<br>62 | (c)<br>80<br>30<br>80<br>100<br>10 |
|                                              |                       |                                |                              | China clay                                                    |                                   | 30                    | 30                                 |

At the present time much sanitary ware is covered with an opaque white enamel 1. The enamel is fired on the raw body, as with hard porcelain; it contains potash, lime, and zine oxide, but neither lead oxide nor boric acid, and is rendered opaque with zinc or tin oxide 2. Alkaline calcareous glazes (Bristol glazes) are used to cover such common stoneware as pots and jars used for holding food or chemicals. Stoneware pipes are glazed by a process known as "salting," in which the goods are heated until the requisite degree of vitrification has been reached, after which a quantity of wet salt is thrown into the furnaces. The vapour of sodium chloride is hydrolysed by the water vapour and the alkali set free reacts with the clay on the surfaces of the pipes to produce a hard, insoluble glaze. The process of salt glazing is most successful with bodies rich in silica, the ratio SiO. M2O2 may vary between 4.6 and 12.5, and in practice it is usually about 8.4

Common red and brown pottery ware may be glazed by coating the unfined bodies with powdered lithage, red lead, or galena, and then firing. A brilliant lead glaze is thus produced and its durability is increased if flint and ballclay are also constituents of the glaze

Decoration .- Colourng matters may be applied to the decoration of ceramic products in various ways, e.g.

- (i) They may be introduced into the body, which then becomes a coloured body. It may or may not be glazed
- (n.) A coloured body may be used to cover another body, either for the sake of economy or for decorating the surface with paintings and with reliefs in different colours ("barbotine" or slip painting) The coloured body may afterwards be glazed it desired.
- (in ) The colour may be incorporated into the glaze, which becomes a coloured glaze.
- (iv.) Colours may be affixed to the surface of the body and the whole then glazed (under-ylaze colouring).
- (v) The colour may be mixed with a flux, ie made into a vitifiable colour and applied to the surface of the glaze (over-glaze or on-glaze colouring).

<sup>&</sup>lt;sup>1</sup> This is superior to the older yet still wide'y adopted practice of coating the articles with a body made of white burning clay or a mixture similar to an earthenware body and then applying the glaze to this fine coating.

<sup>2</sup> See Purdy, Trans. Amer. Cer. Soc., 1902, 4, 61; 1903, 5, 136.

<sup>3</sup> See Barringer, Trans. Amer. Cer. Soc., 1902, 4, 211.

Two or more of these processes may be utilised in decorating the same object.

The best examples of coloured bodies are to be found in coloured parians, porcelains and jaspeis, mosaic tiles and tessera. The bodies are rich in felspar, in which the colouring matters dissolve. The percentages of certain colouring matters that may be added to the body mixtures to produce various colours are as follows.

```
Pal. blue 2 per cent, of cobalt oxide and 5 per cent of zinc oxide
Strong blue 5 per cent of cobalt oxide
Green 2 per cent of cobalt oxide and 1 per cent, of chrome oxide
Blue-green 2.5 per cent of chromic oxide, 1.75 per cent of cobalt oxide, and
1 25 per cent of zinc oxide
Bronze green 5 per cent of mekel oxide
Bronze 1 to 5 per cent of fierre oxide
Green 1 to 5 per cent of ferric oxide
Flack 7 per cent of ferric oxide
Hack 7 per cent of ferric oxide and 3 per cent, of cobalt oxide
```

Coloured glazes are usually applied to white bothes. In most cases the basis of the glaze is a firt rich in lead. For use with soft porcelains and white earthenwares Salvetat proposed the following recipe for the firt

| 15 1 5 5          |   |   |            |
|-------------------|---|---|------------|
| Red lead<br>Flint |   |   | 2000 parts |
| Calcium borate    | • |   | . 1000 ,   |
| Catemin norate    |   | • | 500 ,,     |

The glaze prepared from these amounts of raw materials may be mixed with various metallic oxides to produce coloured glazes. Thus, blue requires 10 to 125 parts of colait oxide, blue green, 100 to 500 of cupic oxide, wory to deep yellow, 70 to 200 of ferric oxide, madder to purplish-brown, 70 to 125 of manganese dioxide, etc.

From terra-cotta bodies that fire to orange or red tints it is only possible to produce red, brown, and black wares by the use of coloured glazes. For other colours the bodies must either be covered with ships or decorated with enamels. On the other hand, with yellowish terra-cotta bodies there is little difficulty in producing yellow, green, and blue wares.

Colonical terra-cotta bodies may be decorated with *enamels*. With highly calcareous bodies a sodium lead silicate rendered opaque by tin dioxide is used to produce a white enamel, and this is coloured by the addition of suitable materials, e.g. lead antimonate for yellow, cobalt oxide for blue, cupric oxide for green, manganese dioxide for violet, etc.

Under glaze colours are applied to pottery bodies before they are glazed. Accordingly, they must resist the action of the glaze at its softening point and be sufficiently infusible to remain still under it, moreover, they must have such rates of expansion as to agree with both body and glaze, since otherwise the glaze will fly off in flakes. The ingredients commonly used in the preparation of under-glaze colours are as follows (Burton):—

```
Black 8 of from chromate, 3 of thinangame tetroxide, 3 of cobalt oxide, and 1 of flint.

Dark blue: 4 of cobalt oxide, 1 of chalk, and 1 of flint.

Azure blue: 3 of cobalt oxide and 60 of animonous alum.

Blue-green: 12 of zinc oxide, 24 of chromic oxide, 4 of cobalt oxide, 12 of chalk, and 12 of boax.

Brown: 8 of from chromate, 4 of rinc oxide, and 2 of ferric oxide.

Reddish-brown: 6 of precipitated from chromate, 3 of lithange, and 20 of zinc oxide.

Fink: 100 of stannic oxide, 34 of chalk, 1 of chromic oxide, and 5 of silica

Fellow: 6 of red lead, 3 of antimony oxide, 1 of stannic oxide, and 2 of flint.
```

Each of the preceding colours is prepared by finely grinding and thoroughly mixing the constituents, strongly calcining the mixture, and regunding the product.

On-glaze colours are fusible coloured glasses which, when applied to an article already glazed and then refired, become fused to the surface. The basis of an on-glaze colour is a readily fusible glass (termed a flux) prepared by fusing together red lead, silica, and borax in suitable proportions. The following compositions are largely used in Staffordshire (Burton):—

|          | 1      | 2    | 3    | 4    | 5    |
|----------|--------|------|------|------|------|
| Red lead | . 64 5 | 50 0 | 30 5 | 15·8 | 40°0 |
|          | 21 5   | 16 7 | 41 7 | 21·2 | 20°0 |
|          | 14 0   | 33 3 | 27 8 | 63·0 | 40°0 |

The powdered flux is mixed with suitable colouring materials, and then, in general, the mixture is fused, cooled, and re-ground to powder. The following compositions are given by Burton (fluxes as in preceding table) --

Pale yellow • 11 of potassium antimonate, 6 of 2mc carbonate, and 80 of flux No. 1.
Orange 14 of potassium antimonate, 6 of ferric oxide, and 80 of flux No. 1.
Uranium orange 25 of yellow oxide of manium and 75 of flux No. 1.
Pale ppg 5 of cobalt carbonate, 3 of ferric hydroxide, and 92 of flux No. 2.
Strong given 6 of cobalt carbonate, 13 of ferric hydroxide, and 80 of flux No. 2.
Strong black 10 of cobalt carbonate, 10 of ferric hydroxide, and 80 of flux No. 2.
Iridium black 25 of iridium seequi oxide and 75 of flux No. 2.
Uranium black 25 of black oxide of manium and 75 of flux No. 2.
Strong blace 13 of cobalt carbonate, 14 of zinc oxide, and 80 of flux No. 2.
Strong blace 13 of cobalt carbonate, 25 of zinc carbonate, and 61 of flux No. 2
Sea green 20 of copper carbonate and 80 of flux No. 2.
Vellow ich brown 10 of ferric oxide, 15 of zinc carbonate, and 75 of flux No. 2
Red 25 of ferric oxide and 75 of flux No. 2 1
Rose 12 of Purple of Cassius, 3 of gold oxide, and 90 of flux No. 4 if Blansk-green 6 of cobalt carbonate, 19 of chromic oxide, and 75 of flux No. 5

Both under-glaze and on-glaze colours are mixed with resmons media and applied by painting, stencilling, transferring from printed or hthographed patterns, or in a number of other ways.

Decoration with metals is often applied to pottery. Gold, platinum, and silver are used for this purpose, usually for over glaze decoration. Gold, platinum, silver, copper, iron, or lead may also be applied to pottery in such a manner as to cover it with a very thin, iridescent, metallic film (lustic pottery).

 $<sup>^1</sup>$  For the production of ferric oxide red, ferrous sulphate is calcined at 400°-420° until a brilliant red oxide is obtained , the oxide is then simply mixed with the flux and not fused with it

with it

The colours obtained from gold are carefully heated to suitable temperatures below a
red heat

## CHAPTER V.

# ULTRAMARINE.1

It has been already mentioned (p 95) that the blue mineral lazurite, extracted from Lipis Lizuli took by powdering and washing with water, was for many years prized as a valuable pigment under the name of ultramarine. The composition of the natural substance was determined by Klaproth 2 and by Clement and Desormes, and m 1828 J B Guinet commenced the manufacture of artificial ultramarine. The artificial product is now, and has been for some years, produced on a very large scale, the annual production ranging from ten to fitteen thousand tons !

Manufacture of Ultramarine Blue.- The constituent elements of ultramatine are sodium, aluminium, silicon, sulphur, and oxygen pigment is prepared by a method which may be looked upon as the calculation of a mixture of an aluminosthere acid and sodium sulphide. The raw materials are as follows

- (i) Clay. China clay or high grade pottery clays are used.
- Kieselgulu is usually employed, but occasionally finely (u.) Silica powdered quartz is used
- (iii) Glauber salt. The best quality of sodium sulphate, free from iron or acid, is calcuid and finely powdered

  (iv) Sodium carbonate The best commercial soda ash is calcuid and
- (v ) Sulphui Rod or stick sulphur practically free from non-volatile matter is used
- Pine charcoal is usually preferred, it is replaced at times (vi) Carbon by pitch, tar. colophony, etc

<sup>1</sup> For further information on ultramitme, see Zerr and Rubeneamp, Treatise on Colour Manutacture, transl. by Mayer (C. Griffin & Co., Ltd., 1999). Bersch, Mineral and Lake Pigmonts, transl. by Wright (Scott, Greenwood & Son, 1901). Parry and Coste, The Thermstry of Tennents (Scott, Greenwood & Son, 1902). Floopp. Dictionary of Applied Chemistry (Longmans & Co., 1912-1913). vol. v., Morsan, Traite de chimie minerale (Paris, 1905), wol. v., Dammer, Hondbanh der anorquineschen Chemier Stuttgart, 1893), vol. ur., R. Hoftworm, Dist Ultramation (Frankfurt, 1873, and Brimswick, 1902). Friend, An Introduction to the Chemistry of Parists (Longmans & Co., 1910). 2 Klapnoth, Jan. Chim., 1797, 21, 150. 3 Cl. ment and Desormes, ibid., 1806, 57, 317. Other analyses are cited by Guckelberger, Annalen, 1882, 213, 208.

<sup>&</sup>lt;sup>2</sup> Ch ment and Desonnes, ibid., 1806, 57, 317 Other analyses are cited by Guckelberger,
Annih, 1882, 213, 208

<sup>4</sup> On the Instant of artificial ulti uname, see Lon, Mem Acad Sci. Lyons, 1878, 23,
333, Heintze, I makt Chem, 1891, [n], 43, 98, Morssan, Tranté de chemu menérale (1905),
vol v. p. 114. See also the following early memoris: Vanquelin, Ann. Chem, 1814, 89, 88;
kublimann, Ann. Chem, Phys., 1829, 40, 139, J. B. Guinet, ibid., 1831, 46, 433. Mérimée,
Breunlin, Annalen, 1856, 97, 295, Ann. Chem, Phys., 1856, [m], 48, 64.

Three chief varieties of ultramarine blue are found in commerce, viz.:-(i) Glauber salt or sulphate ultramarine, the palest variety, which

possesses small covering power

(ii) Soila ultramarine poor in sulphur, a pure blue variety, darker and possessed of more covering power than (1)

(ni.) Soda ultramarine rich in sulphur and silica, the darkest variety. It possesses considerable "body" and is more resistant towards alum than (1.)

or (11).

The first step in the manufacture consists in the powdering and mixing of the ingredients. The most suitable proportions are given by Zerr and Rubencamp as follows --

| Ultramarine Blue    | Pale, or (1) | Pale, or (1) Medium, or (11) |     |  |  |
|---------------------|--------------|------------------------------|-----|--|--|
|                     | -            | ,                            |     |  |  |
| China clay          | 100          | 100                          | 100 |  |  |
| Soda                | 9            | 100                          | 103 |  |  |
| Glauber salt        | 120          | . 0                          | 0   |  |  |
| Carbon .            | 25           | 1 12                         | 4   |  |  |
| Silica (Kieselguhi) | ; 0          | 0                            | 16  |  |  |
| Sulphui             | 16           | 60                           | 117 |  |  |
| 1                   |              | 1                            | i   |  |  |

The second step consists in roasting the mixed materials. This is carried out either by (a) the direct or (b) the indirect process. All ultramarines rich in silica and a certain amount of the other varieties poorer in silica are made by the former method, but the greater part of the ultramarme poor m sihea is prepared by the latter method, in which ultramarine green is formed as an intermediate product. Both these are dry processes, wet methods have been described and patented, but do not seem to have been adopted on a large scale 1

In the direct process the roasting is carried out in such a manner that air has access to the material. Charges amounting to as much as 5 tons may be treated by this process, the heating being carried out in stone troughs, 20 feet long and 10 feet wide, placed in a furnace in such a manner that they can be reached on all sides by the flames The troughs are filled to a depth of 12 to 16 mches, covered with firebricks, and heated for about three weeks 2 It has been proposed to hasten the preparation by adding sodium chlorate as an oxidising agent 3

In the induced process the material is filled into crucibles, the lids are fixed on with a mortar of clay, the crueibles stacked in layers in a furnace and heated, slowly at first, eventually to a bright red heat. By this means air is oxcluded from the mass and green ultramarine produced. The cruebles are opened when the furnace has completely cooled down, and the green product is removed and powdered. It is then converted into blue ultra-marine by roasting it with sulphur. The process is carried out in non-or-fireclay retorts, either (1) by introducing the given ultramarine, together with the

Knapp, J. prakt. Chem., 1885, [n], 32, 375, 1888, [n], 38, 48; M'Ivol, Eng. Pat.,
 9200 (1880): M'Ivor and Cluickshank, Eng. Pat., 19,411 (1892).
 Numerous special furnaces have been patented. Eng. Pat., 18,527 (1890); Fr. Pat.,
 399,211; 400,103; 403,247; 407,089 (1908), 410,055 (1909); 425,585 (1910).
 Fr. Pat., 391,779. 391,780 (1908).

requisite amount of sulphur, 1 into a red-hot retort and raking the mass over at intervals until the sulphur has burnt away, or (n) by heating the green ultramarme and gradually adding the sulphur, each portion of which is allowed to burn away before the next is added

The crude ultrainarme blue, however it has been prepared, is extracted with hot water to remove soluble salts, sodium sulphate being recovered from the washings. The Idue is, while wet, ground to the requisite degree of fineness and separated into particles of different sizes by levigation. The various grades are then dired in sheds, on drying heartles, in special drying rooms, or by any other convenient method. Finally, the discd products are crushed and sifted. The cheaper brands of ultramarme are usually adulterated with 10-50 per cent of gypsum or other cheap filler.<sup>2</sup>

According to Singer 8 ultramarine and analogous bodies may be prepared by treating reolites or similar compounds with sulphides, hydrosulphides, polysulphides or oxysulphides of the alkali or alkaline earth metals

Properties. Ultramarine blue crystallises in the cubic system, isomorphous with the immerals sodalite and hanyinte. The commerced article consists of a bright, azire blue, impalpable powder, insoluble in water, minute particles remain suspended in that liquid for a long time, the whole resembling very closely an inorganic suspension colloid. It withstands a red heat fairly well, but loses some of its brilliancy and turns somewhat greenish in coloni. When heated with water under pressure to 200 - 300° it leaves a colourless regidue free from sulphur, and sodium sulphide passes into solution. It also loses its sulphur when heated with merenic oxide.5 It is stable towards alkalies, but is readily decomposed by dilute nuneral acids with the precipitation of sulphin and the evolution of hydrogen sulphide 5 It is accordingly very remarkable that cold, concentrated (or funning) sulphuric acid, or a mixture of acetic anhydride and glacial acetic acid, has no action upon it 7. Ultramaine possesses hydraulic properties and increases the binding power of cement 8

<sup>1</sup> Seven per cent with soda green and ten per cent with Glinber salt green

<sup>&</sup>lt;sup>1</sup> Seven per cent with sola giern and ten per cent with Glauber salt green <sup>2</sup> For further details concerning the manutor true of ultramarine, see E. Gumet, Ann. Chem Phys., 1878, 1971, 122. Raylins, A See Chem Ind., 1887, 6, 791, Puchot and Granger, thid., 1888, 7, 573, Robing, Chem Ant., 1883, 7, 567, J. Winder, and., 1890, 14, 1119, Jordan, Zeitsch augus Chem., 1893, 6, 684, and the works cited on p. 185. Sugger, D.R.P., No. 221-314 (1909).
<sup>4</sup> Ebell, Ber., 1883, 16, 2129.
<sup>7</sup> Chaburé and Levallors, Compt. rand., 1906, 143, 222.
<sup>6</sup> As a matter of fact, both sulphin droyde and hydrogen sulphide are evolved. The latter, however, is in excess, and part of it is a ed up in decomposing the sulphin droxide. In the presence of an arsente on a cadmium-salt the hydrogen sulphide may be used up in forming arsenous or cadmium sulphide, and the evolution of sulphin droxide may be ned up in forming arsenous or cadmium-shiphede, and the evolution of sulphin droxide may be no bestived. See Guckelberger, Annalen, 1882, 213, 182.
According to 1. Winder (Zeitsch augus Chem., 1912, 77, 200), ultramarine blue losse one-fourth of its sulphin as hydrogen sulphide when treated with dilute hydrochloric acid. This is peculiar, however, to the sample of ultramarine he used. Other samples give

This is peculiar, however, to the sample of idition time be used. Other samples give different results, ear sample 7 quoted in the table on p 133 bost one-third of its sulphur as hydrogen sulphide, and two-thirds of this hydrogen sulphide was used in on decomposing the sulphur dioxide simultaneously produced (B. Campbell, private communication; see also Guckelberger, loc cut

associates, for all 1 of R. A. Hofmann and Metzener, Et 2, 1905, 38 2482; L. Wunder, Zeitsch anora Chem., 1912, 77, 209 When heated with concentrated sulplanne and a white residue is ultimately obtained, the blue codoni group place successively to like black, greenish, black, purple, greenish-grey, and white The purple naticial agrees in its composition and general properties with mole altramarine (B. Campbell private communication)

Rohland, Zeitsch, angew. Chem., 1904, 17, 609.

Ultramarine blue loses part of its sulphur when heated in hydrogen to 450°, but still retains its blue colour When heated to 250° in a stream of chlorine, hydrogen chloride, and steam, it is transformed into wolet ultramavine, 1 which is transformed into redultramarine when heated to  $130^{\circ}$ – $150^{\circ}$  m a stream of hydrogen chloride 2. Blue ultramarine may be directly transformed into the red substance by heating to 400° in nitric oxide.3 The properties of the red and violet ultramarmes have been examined by L. Wunder. Each is produced from the blue by loss of sodium and sulphur; in the conversion from blue into red, half of each of these constituents is removed. Both the red and the violet products are decomposed by hydrochloric acid with the evolution of sulphur dioxide 4 and the separation of sulphur, and both lose a further quantity of sulphur when heated in hydrogen, leaving, however, bright blue residues.

Ultramanne blue is converted into a white solid, with the loss of sodium and sulphur, when heated under pressure to 150° with a solution of phosphorus in earbon tetrachloride. The product loses water and sulphur dioxide when heated, the residue becoming green and then blue. With hydrochloric acid it evolves hydrogen sulphide, and when treated with sodium hypobromite it turns bright blue

When commercial ultramarine blue is heated to 120'-140° under pressure with aqueous silver intrate for fifteen hours, it is converted into a dark yellow silver ultramarine, silver intiite and intric oxide also being produced, but not silver sulphide.6 The silver ultramatine is produced by the replacement of sodium in the blue ultramarine by silver, the complete substitution, however, being difficult to effect. By heating silver ultramarine with molten metallic chlorides it is possible to replace the silver by numerous other metals. The preparation of mercurous, cadmium, lead, lithium, and calcium ultramarines has also been effected directly from ultramarine blue by heating under pressure with aqueous salt solutions.7 The replacement of the sodium by the tervalent metals aluminium, chromium, and iron cannot, however, be thus effected, the ultramarine being decomposed, but L. Wunder considers that aluminium, chromium, and iron ultramarines are produced as intermediate products which are very unstable and decompose It has been known for years that a solution of alum slowly rapidly decolorises ultramarine blue, and the change has been attributed to the hydrolysis of the salt and consequent presence of free acid in the solution. In connection with the various ultramarines of different metals, it is possibly of considerable significance that only those of the alkali metals are blue in

<sup>1</sup> J. Wunder, Ber., 1876, 9, 295; Chem Zett., 1890, 14, 1119, 1906, 30, 61, 78; 1911, 35, 221, L. Wunder, Zeitsch. anorg Chem., 1912, 77, 209; cf. Zeitner, Ber., 1875, 8, 259, 353

2 J. Wunder, loc. cit., cf. Zeitner, loc. cit.; Scheffer, Ber., 1873, 6, 1450; Buchner, Dingl. poly. J., 1879, 231, 446, Ber., 1874, 7, 990, R. Hollmann, Annalea, 1878, 194, 1.

L. Wunder, toc. ett
 Cf. footnote 6 on p. 137.
 L. Wunder, Zeitsch. anorg Chem., 1912, 77, 209, Chem. Zeit., 1913, 37, 1017. For other white ultramaines, see Philipp, Ber., 1876, 9, 1109, 1877, 10, 1227; Annalen, 1876, 182, 132; 1878, 191, 1, Bottinger, Annalen, 1876, 182, 311, R. Hollmann, ibid., 1878,

<sup>194, 1
194, 1
194, 1
196, 1
197, 101,</sup> poly J., 1874, 212, 232, Heumann, Ber., 1877, 10, 991, 1315, 1888;
1879, 12, 60, 784; Annalen, 1879, 199, 253; 1880, 201, 262, 203, 174, Philipp, loc. ct.;
de Forciand and Ballin, Bull. Soc chim., 1878, [11], 30, 112, Chabine and Levallois,
Compt. rend., 1906, 143, 222.
7 L. Wunder, Zeitsch. anorg. Chem., 1912, 77, 209.

colour 1 Thus, the manganese compound is grey, the silver compound is

When silver ultramarine is heated under pressure with a suitable organic rodide, the silver is replaced by an organic radicle. In this manner ethyl, amyl, benzyl, and phonyl oftramarmes have been propared 2. Then formation is not merely a question of substituting silver for an organic radicle, since part of the sulphur is also removed. The organic ultramatimes do not evolve hydrogen sulphide when treated with cold hydrochloric acid. When heated with sodium chloride, however, they give rise to a blue sodium When heated alone, organic sulphides (e.g. ethylsulphide) are ultramarme evolved

Green ultramarine has been already mentioned as an intermediate product in the manufacture of the commercial blue variety. In the conversion of the green into the blue, sodium is removed but sulphur is not added on; the conversion may be effected by heating in a scaled tube with water 3

When ultramarme red is heated out of contact with an it loses a little in weight and is converted into a blue product, like the commercial blue, but unlike the red ultramarine, this new blue product evolves hydrogen sulphide when treated with hydrochloric acid. It differs from the commercial blue in that it is not reduced to a white product by the action of phosphorus, and is not transformed into a violet product by heating to 170° -250 in chloring and hydrogen chloride.

Uses,- Owing to its brilliancy as a body colour and high colouring power, ultramarine blue is largely used in the preparation of blue paints in calico and wall paper printing, for colouring writing paper and printing ink, blueing mottled soap, etc. It is also of value as a whitening agent, coraccting the yellow tint of paper, cotton and linen goods, whitewash, soap, starch, sugar, etc. It also finds considerable application in the manifacture of laundry "blue"

Constitution.—It has been already mentioned (p. 136) that several grades of ultramarine blue are manufactured, differing considerably in their siliea and sulphur content. Numerous analyses of ultramarme blue have been published, a selection is given in the following table  $\sim t$ 

| nent. | 1                | 2    | 1 3       |                                                   |                                                                        |                                                                                                   |                                                                                               |
|-------|------------------|------|-----------|---------------------------------------------------|------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------|
|       |                  |      | 1 "       | 4                                                 | ن ا                                                                    | 6                                                                                                 | 7                                                                                             |
|       | - İ              |      | 1         | i                                                 | -                                                                      |                                                                                                   |                                                                                               |
|       | 40 7             | 37.9 | 424       | 110                                               | 11 3                                                                   | 3 × 3                                                                                             | 10.4                                                                                          |
|       | 24.0             | 30 5 | 24 0      | 211                                               | 24.5                                                                   | 29 2                                                                                              | 23.6                                                                                          |
|       |                  |      |           | ٠.                                                | 1                                                                      |                                                                                                   | 1.1                                                                                           |
|       | 13.6             | 8 4  | 15.4      | 140                                               | 133                                                                    | 43                                                                                                | 13 6                                                                                          |
|       | 17 1             | 16.9 | 13 9      | 16.1                                              | 1115                                                                   | 15.5                                                                                              | 11.6                                                                                          |
|       |                  | i    | 0.7       | i                                                 | 1                                                                      | ł                                                                                                 |                                                                                               |
|       |                  |      | i         | i                                                 | 04                                                                     | 0.7                                                                                               |                                                                                               |
|       | ·<br>·<br>·<br>· | 24.0 | 24.0 30.5 | 24 0 30 5 24 0<br>13 6 8 9 15 4<br>17 1 16 9 13 9 | 24.0 30.5 1 24.0 24.1 24.1 24.0 13.6 8.9 15.4 14.0 17.1 16.9 13.9 16.4 | 24.0 30.5 24.0 24.1 24.5<br>13.6 8.9 15.4 14.0 13.3<br>17.1 16.9 13.9 16.4 11.5<br>10.7 10.7 11.5 | 24.0 30.5 24.0 24.1 24.5 29.2<br>13.6 8.9 15.4 14.0 13.3 9.3<br>17.1 16.9 13.9 16.4 11.5 15.5 |

<sup>1</sup> De Forciand, Bull Soc. chim., 1879, [n ] 31, 161, L. Wunder, lor cut
2 De Forciand, Compt. rend. 1879, 88, 30, Bull Soc chim., 1879, [n ], 31, 161;
Chabrié and Levallois loc cut, et Singer, D.R.P., No. 121,344 (1909).
3 Philipp, Ber. 1870, 9, 110.
4 L. Wunder, Zeitsch amory Chem., 1913, 79, 343
8 Samples 1 and 2 analysed by R. Hodmain (Annalen, 1878, 194, 1), sample 3 by
Parry and Costo (The Chemistry of Francists, 1962), sample 4 is the mean of several analyses by Guckelbergei (Annalen, 1882, 213, 182), samples 5 and 6 by L. Wunder

It is clear that different samples of ultramarine blue may have widely different compositions. According to L. Wunder, the substance always contains a little "water of constitution

Owing to the highly complex chemical constitution of the ultramarines and their derivatives, chemists are not yet agreed as to the chemical formulae to be assigned to them — It is quite possible that the commercial ultra-marines contain a number of closely related compounds in isomorphous mixture. The ultramatures are probably sodium aluminosilicates modified in some manner by the presence of sulphur in the molecules Various suggestions as to the position of the sulphur in an ultramarine molecule have been put forward, thus, Wunder regards the sulphur liberated in the form of hydrogen sulphide by acids as being present in the grouping > Al - S - Na

$$>$$
Al - S - Na + 2HCl =  $>$ AlCl + H<sub>a</sub>S + NaCl,

while groupings such as >Si(ONa)(SONa) lead to deposition of sulphur when the molecule is broken up by acids -

$$>$$
Si $\left\langle \begin{array}{c} 0 \text{ Na} \\ \text{S,0 Na} \end{array} \right. \rightarrow \left. >$ Si $\left\langle \begin{array}{c} 0.11 \\ \text{S,0,11} \end{array} \right. \rightarrow \left. >$ Si $\left. \left. 0 + 11_2 0 + \text{S.} \right. \right.$ 

W. and D. Asch consider that there is an atom of oxygen between the aluminum and the sulphur, but otherwise the corresponding part of their formula is similar to that of Wunder

For further information on the constitution of ultramarme the reader is referred to the hterature !

The Colour of Ultramarine.—The cause of the beautiful colour that ultramarine possesses is at present unknown  $^2$  . Two views are advocated first, that there is a definite chemical individual (or series of alhed individuals) which constitutes the basis of ultramarine and has a blue colour, and secondly, that the colour is more or less an accidental circumstance. On the second view, it is considered that the basis of ultramarine

<sup>[</sup>Zeitsch, anorg Chem., 1912, 77, 200], sample 7 by B. Campbell (private communication). Numerous other analyses are quoted by Guckelberger (Loc. et ), Philipp (.Innalen, 1876, 184, 182; 1878, 191, 1), Bottinger (Lod., 1876, 182, 311), Heumann (Lod., 1879, 199, 253, 1880, 201, 202), Jordan (Zeitsch anew Chem., 1893, 6, 684), and de Forciand (in Moissan's Trauté de chimie, 1905, vol. iv.); see also Dammer, Handhuch der aurganischen Chemie (Stuttgait, 1893), vol. in

1 Breunlin, Annalen, 1856, 97, 295; Wilkens, Lod., 1856, 99, 21. Beckmann, Lod., 1861, 118, 212, Bottinger, Lod., 1876, 182, 311; Philipp, Lod., 1877, 184, 132, 1878, 191, 1; Heumann, Lod., 1879, 199, 253, 1880, 201, 262, 203, 171, R. Holmann, Lod., 1878, 194, 1, Guckelberger, Lod., 1882, 213, 182, Salasu, Lod., 1889, 251, 97, Dolliuss and Goppelstoder, Bull. Soc. Ind. Muth., 1875, 45, 193, 196; Stein, J. prakt. Chem., 1871, ii.], 3, 33; Pheque, Compt. rend., 1877, 85, 570, Bull. Soc. Anim., 1877, 28, 518, 29, 51; Rickmann, Ber., 1878, 11, 2013, de Forcrand. Mem. Acad. Sci. Lyone, 1879, 24, 141; Silber, Ber., 1880, 13, 1854, F. W. Clarke, Amer. Chem. J., 1888, 10, 126; J. Wunder, Chem., 2ett., 1890, 14, 1119, Brogger and Backstoin, Zeitsch. Angew. Chem., 1914, 18, 209; Puchner, Chem. Zentr., 1896, 1, 1051, Rohland, Zeitsch. Angew. Chem., 1914, 77, 209, Book, Zeitsch. angew. Chem., 1914, 77, 209, Book, Zeitsch. angew. Chem., 1915, 28, 117. Most of the Gorgoung are included in W. and D. Asch, The Silicates in Chemistry and Commerce, trausl. by Scale (Constable, 1913).

\*\*The early view that lapis-lazuli owed its blue colour to copper was disproved in 1762 y Margraff, who attributed it to iron. In 1806 Clément and Désormes showed that some ipecimens of lapis-lazuli were free from iron.

is a colourless, transparent solid, which holds in colloidal suspension or solid solution a small amount of another substance which gives rise to the blue colour. This second substance is generally considered to be a blue form of sulphur.1

The arguments used in favour of the colloidal sulphin theory are largely based on analogy, various reactions being known in which it is very probable that colloidal sulphur is produced and in which a blue colon is developed. Thus, potassium thiocyanate 2 turns blue when melted and heated to 400°, and molten sodium chloride or sulphate becomes bright blue when a little sulphur is added and the mass heated until the initial black turbidity vanishes.3 When alkali polysulphides are added to various boiling organic solvents, blue solutions are obtained, which usually lose their colour when cooled 4 Moreover, at very high temperatures sulphur vapour is blue in colour.5

It cannot be considered, however, that the colloid theory is well established. The objection based upon the fact that ultramarine retains its colour at high temperatures is perhaps not very serious, but it is difficult to see how the theory is to explain the existence of red, violet, and green ultramarines. It would necessitate all the colonis being attributed to colloidal sulphur in various states of division, and the assumption that the blue colour of the commercial article is due to its temperature of formation being such as to produce the sulphur in the requisite state of fineness. Silver ultramarine, however, turns blue when heated with sodium chloride at a comparatively low temperature. Again, in the cases of the blue solutions already mentioned, where the existence of colloidal sulphur is considered probable, the blue coloration only makes its appearance in the fluid state, molten, blue potassium thiocyanate becomes white on solidification If, however, in the manufacture of ultramarme, overheating occurs and the mass fuses, the blue colour is destroyed. Ultramarine still retains its blue coloni at - 181°C . A serious objection to the colloid theory is found in the fact that among the numerous substituted. ultramarmes, only those contaming the alkali metals are blue. W. and D. Asch argue in great detail for the view that the ultramarines are definite chemical compounds of a highly complex nature, and that the various colours are produced by changes in the arrangement of the various atoms within the molecule Accordingly to L. Winder, three conditions must be fulfilled for an ultramarme to possess a blue colour (i) an alkalı metal must be present, (n) part of the sulphur must be directly united to the alkali metal, and

ţ. . .

<sup>&</sup>lt;sup>1</sup> See, e.g., Kuapp. I. prokl. Chem., 1888, [n.], 38, 48, Rohland, Zeitsch. angew. Chem., 1901, 17, 609. J. Hollmann, Chem. Zeit., 1910. 34, 821. Abegg, Handhuch der anorgan.; sehen Chemie, vol. in. pt. 1, p. 120 (1906). W. Biltz, eited in Abegg, opins ed. Doelter, Handbuch der Minealchema (Diesden and Lappig), 1911, etc., vol. in. pt. vii., ef. Stein., J. prakt. Chem., 1876, 14, 387. Gentile, Dingl. poly. J. 1856, 140, 223. 141, 165 [1861].
160, 153, Blackmore, J. Noc. Chem. Ind., 1897, 16, 219. E. Guintel, Mem. Acad. Sci., Lange, 1878, 22, 29.

<sup>100, 100 ;</sup> Disconnect of the Lyons, 1878, 23, 29

2 Nollner, Innalen, 1858, 108, 8 ; Ghes, Chem. News, 1901, 83, 61 ; Milbauer, Zertsch. anora Chem., 1904, 42, 433, 1906, 49, 46.

Paterno and Mazzuechelli, Att. R. Acad Lincei, 1907, [v], 16, 1-465

Paterno and Mazzuechelli, los est. For further work on blue sulphur, see Wohler, Annalen, 1853, 86, 373, N. A. ORoll, J. Russ. Phys. Chem. Soc., 1991, 33, 397, 400 1902, 34, 52,

6 B. Campbell (private communication).

7 W. and D. Asch, opus ct.

8 L. Wunder, Zeitsch. anorg. Chem., 1913, 79, 343, Chem. Zeit., 1913, 37, 1017.

(iii.) another part of the sulphur must be present in a lower state of oxidation (i.e. its valency must be less than six). Neither aluminum nor silicic acid is essential, as the former may be replaced by boron, 1 and the latter by a polyboric acid 2

<sup>1</sup> On boson ultramarine, see J. Hoffmann, Zeitsch. unqew Chem, 1906, 19, 1089; 1907, 20, 568; Chem Ind, 1911, 34, 619

2 Selenium and tellurium may replace the sulphin in ultramaine, the substances thus obtained, however, are not blue; Leykauf, Juhresber, Tech., 1876, p. 555, E. Guimet, Ann. Chim. Phys, 1878, [v.], 13, 102, Pheque, Bull Soc chim., 1877, [ii], 28, 518, 1878, [ii.], 29, 522, 30, 51, Morel, ibid., 1877, [ii], 28, 522.



## CHAPTER VI.

## GALLIUM.1

Symbol, Ga Atomic weight, 69.9 (0 = 16).

Occurrence.—Gallium is one of the carest of the elements, occurring widely distributed in nature, but only in very minute quantities? It is a frequent constituent of zinc blendes, the inchest ore, the black blende from Bensberg, contains 16 milligrams of gallium per kilogram. Other blendes relatively rich in gallium are found at Pierrelitte in the Pyrenecs,3 Alston Moor in Cumberland, and Rio Tuerto in Spain 4

Examination of the oxy-hydrogen flame spectra of numerous immerals has shown that gallium is present in a large number of iron ores, particularly in magnetite, clay-nonstone, and black band ore, but not in siderite. When such ores are smelted, gallium passes into the resulting metal, and Middlesbrough pig-non, which contains 0.003 per cent of gallium, is one of the richest sources of the element known  $^5$  Gallium is constantly associated with alummum and chromium in alummous non ores, such as bauxite and kaolin, and accordingly occurs in commercial aluminium, o in one sample of the commercial metal, 0 017 per cent of gallium was found 7

Gallium has been observed in pyrites and manganese ores, and in meteoric non's It also occurs in certain French mineral waters. Gallium exists in the suns atmosphere w

History. On 27th August 1875, Lecoq de Bonsbaudran observed a new violet line, A4170, in the spark spectrum of some material lie had separated from the zine blende of Pietrefitte, further work led to the discovery of another new line,  $\lambda 1031$ , and established the fact that both lines belonged to

<sup>1</sup> See Gallinia," by Lecoq de Borshandian in Wuitz, Dictionnaire de chimie, Supplement, pt. 2, p. 851. Gallinia is identical with the element austrium discovered by Linnemann. See Linnemann, Monotch, 1886, 7, 773, Pribram, ibid., 1900, 21, 118, Lecoq de Borshandian, Compt. vind., 1886, 102, 1136

2 Vernadski, Bull. Mad. Man. Chim. Phys., 1877, [v.], 10, 136

4 Hartley and Ramage, Proc. Roy. See, 1897, 60, 35, 393, Trans. Chim. Soc., 1897, 71, 513. See also Urbain, Compt. vend., 1909, 149, 602, Angel del Campo y Cerdán, Anal. Fis. Quim., 1914, 12, 80

<sup>533</sup> See also Urbam, Compt. rend., 1909, 149, 602, Angel del Campo y e etdan, Ana. Res.

Quim., 1914, 12, 80

Hartley and Ramage, loc cit.

Hartley and Ramage, Trans Chem. Soc., 1897, 71, 547.

Boulanger and Bandet, Compt. rend., 1913, 157, 714

Hartley and Ramage, Sei Proc. Roy. Dubl. Soc., 1898, 8, 703 Sec. also the following papers dealing with the occurrence of gallium. Kurkland. Justr. Assoc. Adv. Sec., 1893, papers dealing with the occurrence of gallium. Kurkland. Justr. Assoc. Adv. Sec., 1893, Chem. J., 1880, 2, 44. Blenden and Bartlett, J. Soc. Chem. J., 1897, Comwall, Amer.

Bardet, Compt. rend., 1913, 157, 221

Bardet, Compt. rend., 1913, 157, 221

Hartley and Ramage, Sci. Proc. Roy. Dubl. Soc., 1898, 7, 1.

the spectrum of a new metallic element, to which the discoverer patriotically gave the name of gallium in honour of his native country. The first researches on gallium 2 were conducted with only a few milligrams of material; but in 1878 Lecoq de Boishandian and Jungfleisch, starting with 2100 kilograms of Bensherg blende, obtained 62 grams of nearly pure gallium.3

The discovery of gallium marked the inauguration of the periodic In putting forward that scheme for the classification of the elements. Mendeleeff was abliged to assume the existence of several elements not known at that period (1871) One of these hypothetical elements, of atomic weight about 68, he called eka-aluminium, and described the properties it should possess, and it was noticed very shortly after the discovery of gallium that gallium closely resembled the hypothetical eka-aluminium in its behaviour. The following tabular statement will serve to illustrate how closely Mendeléeff succeeded in foretelling the properties of gallium :-- 5

#### Eka-aluminium.

## Atomic weight, c. 69

Metal of density 5 9 and low melting-point, not volatile, unaffected by an , should decompose steam at a red heat and dissolve slowly in saids and alkalies

Oxide should have formula El<sub>2</sub>O<sub>3</sub> density 5.5, and dissolve in acids to form salts of the type ElN<sub>3</sub>. The hydroxide should dissolve in acids and alkalies. There should be a tendency towards the formation of basic salts. The sulphate check for these lates.

formation of basic salts. The sulphate should form alims. The sulphate should be precipitated by H<sub>2</sub>S or (NH<sub>4</sub>)S. The anhydrous chloride should be more volatile

than zine chloride.

The element will probably be discovered by spectrum analysis

#### Gallum

Atomic weight, 69 9

Metal of density 5.94, melting at 30.15°, not volatile, unchanged in an , action on steam not known, dissolves slowly in acids and alkalies

Oxide, Ga<sub>2</sub>O<sub>2</sub>, density not known, dissolves in a ids forming salts GaX. The hydroxide dissolves in a bls and alkalies

Salts readily hydrolyse and form basic salts reading nyaring a said. Salts Alums are known. The salphide can be procriptated by H<sub>2</sub>S or (NH<sub>2</sub>)S, but only under special cusumstance. The but only under special cucumstances. The authydrons obloride is more volatile than zine chloride
Was discovered spectroscopically.

Preparation.—Zinc blende containing gallium is dissolved in aqua regia, using a slight excess of blende, in order to leave no free native acid in solution, and the copper, lead, cadmium, mercury, silver, etc., present are for the most part precipitated by the introduction of strips of metallic zinc. The liquid is filtered while hydrogen is still being evolved, and the filtrate boiled for some hours with a large excess of zinc. The abundant white precipitate which forms, containing alumina, basic salts of zinc, gallium, iron, and schromium, and a little silica, is dissolved in hydrochloric acid, and the preceding processes are repeated on the solution b. The hydrochloric acid solution of the final precipitate is saturated with hydrogen sulphide and

<sup>. 1</sup> Lecoq de Borsbaudran, Compt. rend., 1875, 81, 493.
2 Lecoq de Borsbaudran, Compt. rend., 1875, 81, 1100, 1876, 82, 168, 1036; 1876, 83, 261, 824, 1044; Ann. Chem. Phys., 1877, [v. ], 10, 100.
3 Lecoq de Borsbaudran and Junglersch, Compt. rend., 1878, 86, 475
4 Mendeléell, Compt. rend., 1875, 81, 969
5 Mendeléell, Compt. rend., 1875, 81, 969
6 Mendeléell, loc. cit., and J. Russ. Chem. Soc., 1869, 1, 60; 1871, 3, 47, Annalen

Suppl., 1872, 8, 133.

By proceeding thus far with 10 grams of blende, concentrating the chloride solution and examining its spark spectrum, a qualitative test for gallium may be made.

filtered. From the filtrate, the rine is precipitated as sulplinde after the addition of ammonium acctate and acctic acid. This precipitate contains all the gallium (see p 149), the precipitation is effected in fractions, the filtrate after each operation being spectroscopically examined for gallium. The washed sulphides are converted into chlorides and the cold liquid fractionally precipitated with sodium carbonate, gallium quickly concentrating in the first fractions of the precipitate The precipitates containing gallium are dissolved in sulphuric acid, and the solution gently heated until nearly all the excess of acid has been expelled. The residue is dissolved in cold water, the solution largely diluted, and beiled for some hours - Basic gallium sulphate separates out and is filtered and washed with hot water precipitate is dissolved in sulphuric acid, and any non-present is precipitated by a luge excess of potassium hydroxide. From the filtered solution the gallium is precipitated by a prolonged current of carbon dioxide. The procipitate is dissolved in the minimum amount of sulphinic acid, the solution treated successively with animomium acetate and hydrogen sulphide, and filtered. The gallium is again precipitated as a basic sulphate by Loiling the largely diluted filtrate, the washed precipitate dissolved in sulphuric acid, and an excess of potassium hydroxide added. The liftered solution is then electrolysed to deposit the gallium, using a large platinum cathode. Gallium may also be deposited electrolytically from an aumoniacal solution of the sulphate. The metal, which easily separates from the electrode by bending it under water, is allowed to stand for several hours in very dilute hydrochloric acid, then in dilute potassimi hydroxide at 50 to 60°, and is finally washed

Properties. Gallium, the eka aluminium of Mendeléeff, is a hard, brittle, grey metal having a greenish blue reflex. It melts at the remarkably low temperature of 30 15 (° to a silver-white liquid which is only slightly volatile even at a red heat? In the absence of the solid physe, liquid gallinm possesses to an extraordinary degree the property of remaining in a superfused state. The metal crystallises with great readiness in the tetragonal (or possibly the monoclinic) system, the crystals possessing an octahedral liabit and usually possessing slightly convex faces. The density of the solid is 5.96 at 24.7° 2. Gallium, therefore, like water, expands on freezing. The mean specific heat of the solid metal is 0.079 between 12° and 23°, that of the biquid is 0.0802 between 106° and 119° The latent heat of fusion is 19 03 calories per grain of metal <sup>3</sup> Tho specific resistance exceeds that of the alkali metals <sup>4</sup> Liquid gallium is electronegative to the solid element? The atomic refraction of gallium in its compounds is 14.8 (for the Ha line , Gladstone and Dale's formula). Gallium is

<sup>&</sup>lt;sup>1</sup> For further details and alternative methods, see Lecoq de Bossbandian, Ann Chim, Phys., 1877, [v.], 10, 129, Compt. rend., 1876, 82, 1098, 83, 636, 1881, 93, 815. Wurtz, Dictionnaire de chimie, Supplement, pt. 2, p. 851; Lecoq de Bossbandian and Jungflosch, loc. etc.

Location of Grand, Compt., and . 1876, 83, 611, 1011; Ann. Chim. Phys., 1877, [v | 10, 100.

Bethelot, Compt. rend., 1878, 86, 786

<sup>Betthelot, Compt. rend. 1878, 86, 786
Guntz and Bromewski, Compt. rend. 1808, 147, 1474, J. Chem. phys., 1909, 7, 464.
Regnandl, Compt. rend, 1878, 86, 1477
Gladstone, Proc. Roy. Soc., 1897, 60, 140.
Owen, Proc. K. Akad. Wetensch. Amsterdam, 1911, 14, 637, Ann. Physik, 1912, 127, 657</sup> [1v.], 37, 657

The spark spectrum of gallium is characterised by two violet lines, \$\lambda4172-2\$ and \$\lambda 4033.2, the former being the more intense.\frac{1}{2}

Gallium is only superficially oxidised when heated, even to redness, in air or oxygen. It does not decompose water at 100°, and under an-free water it remains bright and unaffected for a long time. It dissolves slowly in immeral acids. Solid gallium dissolves much faster in hydrochloric acid than does the liquid metal. Warm nitric acid dissolves it slowly, and the best acid solvent is aqua regia. Gallium dissolves in potassium hydroxide, with the liberation of hydrogen. It unites directly with chlorine, bromme, and iodine, in decreasing order of activity 2

Gallium forms two classes of compounds, of the types GaX<sub>2</sub> and GaX<sub>3</sub> (X denoting a univalent acid radicle) The latter closely resemble the compounds of aluminium. Concerning the former, which instantly reduce potassium permanganate in dilute acid solution, very little is known

Atomic and Molecular Weights. -The vapour densities, referred to air as unity, of the two chlorides of gallium have been determined to be 4.82 (at 1000° to 1100°) and 6.13 (from 440° to 606°), corresponding to the molecular weights 140 and 178 respectively  $(O_2=32)^{3}$ . Since the chlorides contain 51 0 per cent, and 40 0 per cent of gallium respectively, the weights of metal in the preceding molecular weights are 714 and 712. Hence Avogadro's hypothesis leads to the approximate value 71 for the atomic weight of gallium, and to the molecular formulae GaCl, and GaCl, for the

The specific heat of gallium (p. 115) supports this view, since it indicates an atomic weight of about 80, and it is confirmed by the fact that gallium sulphate forms a series of double sulphates, isomorphous with ordinary alum, Al<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub> K<sub>2</sub>SO<sub>4</sub> 24H<sub>2</sub>O, which double salts, on the assumption that the approximate atomic weight of gallium is 70, must be formulated

 $Ga_2(SO_4)_3$   $R_2^2SO_4$ :24H $_2O_5$  where R=K,  $Rb_2^2C_5$ ,  $NH_{10}$  or TH. The preceding approximate value for the atomic weight of gallium indicates that it is three times the combining weight of the metal in its oxide and highest halogen compounds. The value at present accepted for the atomic weight, Ga = 69.9, results upon a single analysis of gallium ammonium alum and a single synthesis of gallium sesqui-oxide s

The atomic weight of gallium was calculated by Locoq de Borsbaudran, before sufficient material was available for an experimental determination, from considerations based upon a comparison of the wave-lengths of the spectrum lines of gallium and of other alhed elements of known atomic weights. His method of calculation leads to the value Ga = 69.86 %

The molecule of gallium is monatomic in dilute solution in mercury.

<sup>1</sup> Lecoq de Boisbaudian, Compt. rend., 1876, 82, 168, 1887, 104, 1581, 1892, 114, 315, Livening and Dewar, Proc. Roy. Soc., 1879, 28, 471; Kayser, Hambuch der Spellenstopte (Leipzig, 1900-12), vol. v. p. 460, Hariley and Moss, Proc. Roy. Soc., 1912, A, 87, 38. On series in the spectrum of gallium, see p. 3. On the high-frequency spectrum, see de Broghe, Compt. rend., 1914, 159, 304.

2 Lecoq de Boisbaudran, Compt. rend., 1876, 83, 663, 824.

3 Nilson and Pottersson, Trans. Chem. Soc., 1888, 53, 811.

4 Lecoq de Boisbaudran, Compt. rend., 1881, 93, 294, 329.

5 Lecoq de Boisbaudran, Compt. rend., 1878, 86, 941, see F. W. Clarke, A. Readculation of the Atomic Weights ("Smithsoman Miscellaficous Collections," vol. 51, No. 3), 3rd ed., 1910, p. 269.

 <sup>1910,</sup> p. 269.
 See the article "Gallium" in Wuitt, Dictionnaire de chimic, Suppliment, pt. 2, p. 859;
 Miss Freund, The Study of Chemical Composition (Cambridge University Press, 1904), p. 491. 7 Ramsay, Trans. Chem Soc., 1889, 55, 521.

### COMPOUNDS OF GALLIUM.

Alloys—Galhum easily alloys with aluminum, the inixtures rich in galhum being liquid at ordinary temperatures. The alloys readily decompose water, liberating hydrogen and leaving a solid residue of metallic galhum and aluminum hydroxide.1 Gallinia also alloys readily with indimin

Gallium dichloride, GaCl, prepared by heating the trichloride with excess of gallium and distilling the product in carbon dioxide, forms white, transparent crystals, melting at 164° and boiling at 535°. The liquid exhibits the phenomenon of superlusion to a remarkable degree, as, indeed, do all the halogen compounds of gallium. The vapour density at 1000° to 1100' is 4.82 (an - 1), the formula GaCl, corresponding to 4.86. At higher temperatures dissociation takes place, probably into GaCl and chlorine <sup>4</sup>

in most an, the dichloride deliquesces to a clear liquid. The addition of an excess of water leads to precipitation of an oxychloride, hydrogen (and perhaps gallium hydride) being evolved 5

Gallium trichloride, GaClm obtained by heating gallium in chlorine or hydrogen chloride, forms long, white needles, melting at 75.5° and boiling at 215° to 220° 6° The vapour has a density of 6.13 (air 1) between 140° and 606°, the simple formula GaCl<sub>3</sub> corresponding to 608° At 1000° the vapour density indicates that dissociation has occurred, while at low temperatures the results obtained by the method of Dumas point to the existence, over the range 237° to 378°, of gaseous Ga<sub>5</sub>Cl<sub>o</sub>, the molecules of which gradually undergo dissociation into GaCl, with further rise of temperature

Molten gallium trichloride has a density of 2°36 at 80°,5 readily absorbs gases, but evolves them on solidification. The crystalline chloride is very deliquescent and dissolves in water with the evolution of considerable heat, producing a colourless solution from which oxychlorides are slowly precipitated." The aqueous solution loves hydrochloric acid when evaporated, leaving an amorphous residue which absorbs water and becomes gelatinous. An acid solution of the trichloride is obtained by dissolving gallium in aqua regia

The bromides of gallium 10 closely resemble the chlorides They are, however, less fusible and less volatile, and do not seem to have been analysed. The iodides of gallium, like the broundes, require further study. Two

appear to exist, a colourless tri-iodide and a yellow di iodide.

Gallium sesqui-oxide, Ga,O, which may be prepared by igniting the hydroxide or intrate, is a white mass, infusible at a red b at - Its specific heat (0° to 100°) is 0 1062  $^{11}$ . The strongly ignited oxide resembles ignited alimning m its resistance to acids and alkahes, and requires to be fused with potassium

Lecoq de Borsbaudian, Compt. rend., 1878, 86, 1201.
 Lecoq de Borsbaudian, thid., 1885, 100, 701.
 Lecoq de Borsbaudian, thid., 1881, 93, 294.
 Nilson and l'etterson, Trans. Chem. Soc., 1888, 53, 814.
 Lecoq de Borsbaudian, Compt. rend., 1881, 93, 294. 1882, 95, 18.
 Lecoq de Borsbaudian, thid., 1881, 93, 329.
 Nilson and Petterson, Trans. Chem. Soc., 1888, 53, 814., Friedel and Ciafts, Compt. 1888, 107, 306.

<sup>\*</sup> Nilson and Pettersson, 17403 Co. m., 500, 300, 300, 1888, 107, 306

\* Lecoq de Bushaudran, Comp' rind., 1881, 93, 294.

"Lecoq de Bushaudran, ibid., 1882, 94, 695

10 Lecoq de Bushaudran, ibid., 1878, 86, 756. with Jungfleisch, ibid., 1878, 86, 577

11 Nilson and Pettersson, Compt. rend., 1880, 91, 232, Ber., 1880, 13, 1459.

hydrogen sulphate in order to be brought into solution. At a red heat, hydrogen reduces the sesqui oxide to a greyish-blue mass, probably a suboxide (GaO?), which dissolves in dilute mineral acids to form solutions which instantly reduce potassium permanganate.\(^1\) At a bright red heat, reduction to the metal can be effected. Gallium sesquioxide is also reduced to gallium when heated with magnesium.2

The oxides of samatium,  $Z_a$  (dysprosium),  $Z_{\beta}$  (terbium), and chromium act as phosphorogens when diluted with gallium sesqui oxide and submitted to the action of cathode rays. The phosphorescence is a beautiful red when

the quantity of chromium sesqui ovide equals 0.1 per cent 3

Gallium hydroxide, Ga(Oll); (1), does not seem to have been analysed. It is obtained as a white precipitate, readily soluble in potassium hydroxide, by adding ammonia to a solution of a gallium salt. Tartarie acid hinders the precipitation

Gallium sulphide, Ga,S3(!), undoubtedly exists, but it has not been obtained pure, since its precipitation can only be effected in the presence of

another insoluble sulphide 1

**Gallium sulphate,**  $Ga_2(SO_4)$ , is a white, crystalline salt. Its specific heat (0° to 100°) is 0.116 °. It is very soluble in water, soluble in 60 per cent. alcohol, but insoluble in other  $^6$ 

Gallium ammonium alum, Ga<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub> (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> 24H<sub>2</sub>O, erystallises from a solution of mixed gallium and ammonium sulphates in combinations of the cube and regular octahedron The aqueous solution precipitates basic salts when heated

The corresponding potassium-, rubidium-, cæsium-, and thallium-gallium alums have been prepared by Soret, who gives the following values for the densities and refractive indices  $(\mu)$  for the D line at ordinary temperature .-

Gallium Silicotungstate.—Gallium silicotungstate is very soluble in Three hydrates are known the first, which has the formula  $2 Ga_{\rm s} O_{\rm q}.3 (SiO_{\rm s} 12 {\rm WO}_{\rm s})$  93 H<sub>2</sub>O, crystallises in regular octahedra . the second has 8711,0 and crystallises in Thombohedra (a c-1 26316), while the third has  $60 H_2 O$  and crystallises in the monoclinic (4) system (a b c= 0.9057 1  $\pm 1.1585$ ,  $\beta = 74^{\circ}$  20'). The first two salts therefore correspond to those of alummum (p. 87) 9

Gallium nitrate, Ga(NO<sub>3</sub>)<sub>4</sub>, Al<sub>2</sub>O, is a white, dehiquescent salt which is completely decomposed at 200°. <sup>10</sup>

Dupné, Compt rend., 1878, 86, 720.
 Winkler, Her., 1890, 23, 772.
 Lecoq de Borsbautdran, Compt. end., 1887, 104, 330, 1581, 105, 784, 1228.
 See p. 149, and Lecoq de Borsbautdran, Compt. rend., 1881, 93, 815.
 Nilson and Petterson, Compt. rend., 1880, 91, 232; Ber., 1880, 13, 1159.
 Lecoq de Borsbaudran, Ann. Chem. Phys., 1877, [v], 10, 126.
 Lecoq de Borsbaudran, Compt. rend., 1875, 81, 1100, 1876, 83, 821; Ann. Chem. 1877, [v], 10, 126.

Phys., 1877, [v.], 10, 128.

Soret, Arch. Sci. phys. nat., 1885 [m], 14, 96, 1898, [m.], 20, 520

Wyrouboff, Bull. Noc franc. Min., 1896, 19, 219

The hydrate with 60H<sub>2</sub>O is possibly triclinic, see Wyrouboff, that, 1905, 28, 237.

Dupić, Compt. rend., 1878, 86, 720.

# DETECTION AND ESTIMATION OF GALLIUM

A solution of a pure gallium salt is not precipitated by hydrogen sulphide m acid, neutral, or alkaline solution, but, in the presence of excess of another element capable of forming an insoluble sulphide, gallium sulphide is carried down completely with the other sulphide in either acetic acid or animoniacal solution Arsemous sulphide is usually best to employ, but the sulphides of silver, of zinc, and particularly of manganese may be used In solutions containing much mineral acid, gallium sulplinde is not precipitated, a fact that is utilised in effecting the separation of gallium from the excess of either arsenic or silver

Ammonium hydroxide or carbonate, and carbonates of the alkah and alkaline earth metals, precipitate gallinin as hydroxide from solutions of its salts The precipitate dissolves appreciably in excess of the soluble precipitants, but precipitation by ammonia is practically complete if excess of ammonia is removed by bothing

Gallium may be precipitated as lerrocyanide, but not as lerricyanide, in solutions containing 33 per cent by volume of concentrated hydrochloric acid, the reaction being extremely delicate. It is completely precipitated as hydroxide by boiling the sulphate or chloride solution with excess of piccipitated cupie hydroxide In the presence of non it is preferable to boil with copper to reduce ferrie to ferrous salt, and then to precipitate the gallium by the addition of precipitated cuprous oxide—The detection of gallium spectro scopically in minerals has already been described (p. 111, footnote 6)

Gallium is weighed as the sesqui oxide? ----

<sup>&</sup>lt;sup>1</sup> Lacoq de Borshaudran, Ann. Ch.m. Phys. 1881, [vi ], 2, 176, 129, Compt. rend., 1881, 93, 815., 1882, 94, 1151, 1227, 1439, 1625, 95, 157, 410, 503, 703, 1192, 1339, 1883, 96, 152, 1696, 1838, 97, 66, 142, 295, 521, 623, 730, 1463, 1884, 98, 741, 781, 99, 526

## CHAPTER VII.

#### INDIUM.

Symbol, In. Atomic weight, 114.8 (0 = 16).

Occurrence. - Indium is one of the rarest of the elements, occurring in minute quantities tairly widely distributed in nature 1. It exists in most zinc blendes,2 in some tungsten ores, and in many specimens of pyrites. It is a constant constituent of tin ores, and occurs in siderites, associated with manganese. Indium has also been observed in some Italian galenas and in other minerals 5

History. -In 1863, Reich and Richter observed, in the spark spectrum of a specimen of Freiberg zinc blende, two new indigo-blue lines. They attributed them to a new metallic element, which they succeeded in isolating Subsequently, the new metal, which had been appropriately named indium, was studied in detail by Winkler.7

Preparation.--Indium is best extracted from sine that has been prepared from blende containing indium. The sine, which may contain 0.1 per cent of indium, is treated with a slight deficit of dilute sulphuric or hydrochloric acid, and the solution allowed to stand in contact with the excess of zinc for several days. From the spongy residue, which contains indium, lead, copper, iron, cadmium, arsenic, and the excess of zinc, pure indium oxide is best prepared by the method due to Bayer.8 The washed residue is dissolved in nitric acid and the solution evaporated with a slight excess of sulphune acid. Iron and indium hydroxides are precipitated by the addition of ammoma to the filtered solution of the sulphates and the washed precipitate dissolved in hydrochloric acid. The nearly neutral solution when boiled with excess of sodium bisulphite yields basic indium sulphite as a fine, crystalline powder. The precipitate is dissolved in aqueous sulphirous acid and the filtered solution heated to boiling, when pure basic indium sulphite is obtained.9

Vernadski, Bull Acad Sci Petrograd, 1910, p. 1129, 1911, p. 187
 Reich and Richter, vide infra, Kachler, J. prakt. Chem., 1865, 96, 447, Cornwall, Chem. News, 1873, 28, 28, Urbain, Compt. rend., 1909, 149, 602, Angel del Campo y

Chem. News, 1043, 20, 20, Urbain, Compc. rena., 1905, 149, 602, Ruger der Campo y Cerdán, Anal. Fis Quim 1914, 12, 80.

3 Hoppe-Seyler, Annalen, 1866, 140, 247; Atkinson, J. Amer. Chem. Soc., 1898, 20, 811.

4 Haitley and Ramago, Trans Chem. Soc., 1897, 71, 533.

5 Do Negri, Ber., 1878, 11, 1249, Gazzetta, 1878, 8, 120; Tanner, Chem. News, 1874,

<sup>Do Negri, Ber., 1818, 11, 1249, transacte, 10,0,0,12, 12, 130, 141.
Reich and Richter, J. prakt. Chem, 1863, 89, 441; 1863, 90, 172; 1864, 92, 480;
Richter, Compt rend, 1867, 64, 827.
Winkler, J. prakt Chem., 1865, 94, 1; 95, 414, 1867, 102, 273, condensed account in Ann Chim. Phys., 1868, [iv.], 13, 490.
Bayer, Annalen, 1871, 158, 372. J. Chem. Soc., 1871, 9, 661
Cf Winkler, loo ext.; Bottgey, J. prakt Chem., 1866, 98, 26, R. Meyer, Annalen, 1869, 150, 137. For the extraction of indium from blonde, see Reich and Richter, loc ext., Winkler, loc. ext.; Weselsky, ibid., 1865, 94, 443; Kachler and Schrotter, ibid., 1865, 95, 441; Stolba, Dingl. poly J., 1870, 198, 223.</sup> 

The basic sulphite is dissolved in sulphuric acid, and indium hydroxide thrown down by the addition of ammonia. The precipitate is converted into indium sesqui-oxide by ignition

Indum oxide may be reduced to the metal by heating in hydrogen or by heating with sodium. In the latter case, the brittle alloy of indium and sodium obtained is decomposed by water and the residual indium fused with sodium carbonate.

Traces of non-may be removed from indium by converting it into the trichloride, adding potassium throeyanate to the feebly acid solution, and extracting the terric throeyanate with other  $^2$ . Pure indium trichloride is also obtained by adding pyridine to its alcoholic solution. A double compound of the chloride with pyridine is precipitated, iron and aluminium chlorides remaining in solution.

Indium is easily deposited electrolytically from a solution of the chloride or sulphate, in the presence of pyridine, hydroxylamine, or formic acid, and fractional electrolysis of the sulphate solution affords the best method of obtaining pure indium. The deposited metal is pressed together, washed, dried at 120°, and fused in a charcoal boat in a current of hydrogen.

Properties.—Indum is a soft, ductile, silver white metal, which melts at 155% and is volatile at a red heat. It crystallises, like aluminum, but unlike zine, in the cubic system, heme electrolytically deposited from its sulphate solution in regular octahedra. Its density is 7.277 at 20% its coefficient of expansion, 0.0000459, and its specific heat (0° to 100°) is 0.0570. The atomic refraction of indium in its compounds is 174 (for the H<sub>a</sub> line. Gladstone and Dale's formula) 12. Indium is diamagnetic. 13

The flame, are, and spark spectra of indium are characterised in the visible region by two bulliant indigo blue lines, 1511-55 and 1101-95.11. The most parsistent lines in the spark spectrum of indium and therefore the lines that should be looked for when seeking traces of the element, are (Exper and Haschek's wave-lengths) 1511-55,\* 1101-95,\* 3256-22,\* 3039-16,\* 2941-39, 2890-35, 2710-39, 2306-20, those asterisked being the most sensitive 15.

Indium is unaffected by dry air at ordinary temperatures, but at a red

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    Winkler, Ioc. ett., Boltger, J. prikt Chem., 1869, 107, 39
    Dennis and Geet, J. Amer. Chem., Nov., 1901, 26, 437, Ber., 1904, 37, 961.
    Renz, Iev., 1904, 37, 2110. Dennis and Geer, tec. ett.
    Dennis and Geer, Joe cit.
    Tinel, Iev., 1904, 37, 175, Zet/sch anory Chem., 1904, 39, 119, 40, 250; Mathers, Ber., 1907, 40, 1220, J. Amer. Chem. Sov., 1907, 29, 485.
    Tinel, Ioc cit.
    Tinel, Ioc cit.
    Ditte, Compit. rend., 1871, 72, 858.
    Sachs, Zet/sch. Kryst. Min., 1903, 38, 195.
    Richards and Wilson, Carnegae Institution Publications, 1909, No. 118, p. 13, Zeitsch. physikal Chem., 1910, 72, 129.
    Freau, Compt. rend., 1869, 68, 1125.
    Bunsen, Poug. Annalen, 1870, 141, 28, Thil Mag., 1871, [iv.], 41, 161, Ann. Chim. Phys., 1871, [iv.], 23, 50.
    Gladstone, Proc. Roy. Sov., 1897, 60, 140.
    Gwen, Proc. K. Akad. Wetensch. Amsterdam, 1911, 14, 637; Ann. Thysik, 1912, [iv.], 37, 657.
    Kayser, Handbach der Spektreit opin (Leipzig, 1900-1912), vol. v. p. 581, Exner and Ilaschek, Inc. Spektren. d.: Elemente bet normalem Irrack (Leipzig and Wich, 1911); Schulemann, Zeitsch. was Photochem., 1912, 10, 263 (spaik). On senes in the indium spectrum see p. 3.
    Hartley, Phil. Trans., 1884, 175, 1, 325, de Giannent, Compt. 2cad., 1907, 144, 1101; 1910, 151, 308; 1914, 159, 5, Hartley and Moss, Proc. Rey. Soc., 1912, A, 87, 38.
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heat it burns with a blue flame, producing the sesqui-oxide with the liberation of 10116 cals of heat per gram of metal 1 It unites directly with sulphur and the halogens. It is imaffected by boiling water or potassium hydroxide, but dissolves in immeral acids. The action of intric acid is slow, and ammonia is found among the reduction products of the acid 2. The potential differences between indum and molar, tenth-molar, and hundredth-molar solutions of indium trichloride are 0 094, 0 108, and 0 119 volts respectively at 25°, the metal being negative to the solution. The electrolytic solutionpressure of indium, 102 to 103 atmospheres, places it between non and lead in the electromotive series.8

Atomic and Molecular Weights .- The early workers on melnim regarded that element as a duad, and analogous to zme. This view was based upon the facts that indiam occurs naturally in association with zinc, the two metals are very similar in their analytical reactions, and no sodiumor potassium-indium sulphates could be prepared which crystallised in octahedra and could be formulated as almus on the assumption that indium was tervalent. The bivalency of indimin was, however, not acceptable to Mendeleeff, who could only place indium in his periodic table on the assumption that it was a triad, of atomic weight c 114. In support of the tervalency of indium, both Mendelceff and Lothar Meyer advanced various chemical reasons, 4 and Mendelceft and Bunsen each showed, by determinations of the specific heat of the metal, that the tervalency of indimin was a necessary assumption from the point of view of Dulong and Petit's Law 5 Subsequent work has fully substantiated this assumption. In 1873 Roessler<sup>6</sup> succeeded in preparing indium ammonium alim, and in 1885 Soret 7 prepared the rubidium and casium alims. Most convincing of all, in 1888 Milson and Pottersson's prepared and determined the vapour densities of three chlorides of indium, and showed that the results were in agreement with the molecular formulæ InCl. InCl., and InCl., with In 114 approximately. This value has been contrined by another method, which may be outlined here. The  $\textit{specific opacity}^{\,\alpha}$  of an element for N rays of a definite quality is independent

$$\frac{\mathbf{M}}{\mathbf{I}\mathbf{S}} = \frac{m_1}{\epsilon_1} + \frac{m_2}{\epsilon_2} + \frac{m_3}{\epsilon_3} + \cdots$$

Ditte, Compt. rend., 1871, 72, 858. 73, 108.
 Armstrong and Acworth, Trans. Chem. Son., 1877, in. 81.
 Thiel, loc. cit., measured against decimonal edonal electrode (0.620 volts). Ct.
 Erhand, Wied. Annalen, 1881, 14, 501.
 L. Meyer, Annalen Singh., 1870, 7, 351., Mendelécif, dad., 1872, 8, 133, or Chem.
 News, 1879, 40, 231, etc., 1880, 41, 2, etc.
 Mendelecif, Bull. Acad. Sci. Petrograd, 1870, p. 415., Bunsen, Phil. Mag., 1871, [17].

<sup>41, 161.

&</sup>lt;sup>6</sup> Roessler, J prakt Chem., 1873, [n.], 7, 14

<sup>7</sup> Soret, Arch See, phys. nat., 1885, [m.], 14, 96

<sup>8</sup> Nilson and Petiesson, Trans. Chem. Nov., 1888, 53, 814

<sup>9</sup> Given unit mass of a substance (1 deegs un) in the form of a right cylinder, area of base 1 sq. cm., and given that X-rays full mormally on the base and pass through the cylinder, the triction of the rays absorbed by the cylinder measures the specific operaty of cylinder, the friction of the rays absorbed by the cylinder newsures the specific opening of its material

The equitalent transparency of a substance is the mass (in decignants) of a right cylinder 1 sq. cm. in base which, when X-rays are passed through it parallel to its axis, produces the same absorption as a cylinder of parallin way 75 mm. high and 1 sq. cm. in cross section.

If masses  $m_1, m_2, m_3, \ldots$  of elements of equivalent transparences  $c_1, c_2, c_3, \ldots$ 

If masses  $m_1, m_2, m_3, \ldots$  of elements of equivalent transparences  $e_1, e_2, \ldots$  are present in a total mass M of substance of equivalent transparency E, then, since specific opacity is a strictly additive property,

of its state of aggregation and of the temperature, it is also independent of whether the element is free or in combination, so that the specific opacity of a compound may be calculated from the opacities of its constituent elements. It has been found by Benoist that for X-rays of one definite quality, the specific opacity of an element mercases in a regular manner with the atomic This is perhaps best indicated by plotting the equivalent transparencies of the elements (which are proportional to the reciprocals of the opacities) against the atomic weights. The points he in a smooth curve, such as fig [16], curve [1,2] and the curve is somewhat similar to a hyperbola (fig. 16, curve 111), which is the graphical representation of the connection between specific heat and atomic weight (Dulong and Petit's Law) is, however, a separate curve of transparencies for each quality of N-rays used Thus in fig. 16, curve I refers to tays of medium handness, and curve II to soft rays. Therefore, in seeking the atomic weight of an element by the Viays method, each possible multiple of the combining weight is assumed, the elements which would immediately precede and follow it on the curve noted in each case, and the transparencies of each of these elements, and of the element under investigation, determined for a particular quality of X-rays, or better, for two decidedly different qualities. The results have the great advantages over those based upon specific heats that they are not influenced by temperature or physical state and are applicable to gaseous

The preceding method has been applied to the case of indium, both the free element and its acetylacetonate being used. It was found that both for the medium rays (bg 16, curve I) and the soft rays (curve II) the equivalent transparency of indium is very nearly 140. The results point clearly to the value c 114 for the atomic weight !

The atomic weight of indumers therefore three times its combining weight in its highest halogen compounds and its basic oxide. Early determinations of the atomic weight of indium, made by Reich and Richter, Winkler, and Bunsen, were made almost entirely by a faulty method—the synthesis of the sesqui-oxide - and yielded low results. Clo cly concordant results have since been obtained by Thiel's and Mathers, each of whom analysed the trichloride and tubromide of indining to

```
InCl, 31g(1
               51 173 | 100 000 (Thiel)
   In - 111 968
               51-112 | 100 000 (Mathers)
   =111834
InCl_g=3AgB_I
               62 923 100 000 (Thiel)
   = 111.753
               62:932 100 000 (Mathers)
   =111803
```

See note 9 on page 152

<sup>2</sup> Lithnum, carbon, introgen, and oxygeo also fell on the curve but beyond the limit of A By experimenting with the free elements or the a compounds

4 See Benost, Compl. rend., 1897, 124, 146, 1901, 132-325, 545, 772, J. de physique,

<sup>1901, [</sup>m], 10, 653

<sup>1901, [</sup>in ], 10, 653

b Ruch and Richter, J. prakt Chem., 1861, 92, 190

b Winkler, that, 1865, 94, 1, 1867, 102, 250.

7 Bunsen, Poug. Annalia, 1876, 141, 28

a Thiel, Zeitch anna, Chem., 1901, 40, 280

Matheis, J. Amar. Chem. Soc., 1907, 29, 185, 180, 1907, 40, 1220

10 Clarke, A. Realiulation of the Atomic Winglets ("Smithsoman Miscellaneous Collections," vol., 54, No. 3), 3rd ed., 1910, p. 260

The antecedent data have been changed to the following. Ag=107, 880, Cl=35, 457, Bi=79, 916

The most probable result, according to Clarke, is 114.86, and the international value (1916) is In=II4.8.

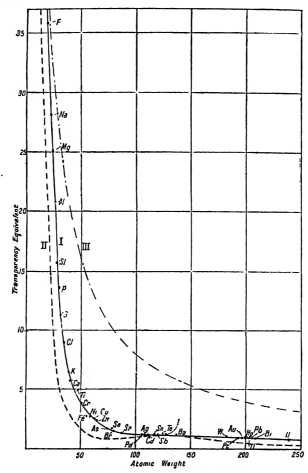


Fig. 16.—Benoist's X-ray transparency curves.

Indium readily dissolves in molten tin, and lowers the freezing-point. The "atomic fall" is 1.86°, the theoretical value for a monatomic molecule being 30°. Hence in dilute solution in tin, indium is mainly diatomic.

<sup>1</sup> Heycock and Neville, Trans Chem. Soc., 1890, 57, 385.

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From the results obtained by Richards and Wilson 1 in a study of the electromotive forces of indium amalgam concentration cells, it follows that indium is essentially monatomic in solution in mercury, or more probably exists as a compound of molecular formula InIIgz, most likely InIIg, 2 Indium is essentially monatomic in dilute solution in sodium, the "atomic fall" being 3 6' and the theoretical value for a monatomic element 4 4°.3

### COMPOUNDS OF INDIUM.

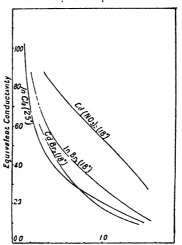
Indum forms derivatives corresponding to the three types InX, InX2, and  $\mathrm{In}\mathrm{X}_{1}\left(\mathrm{X}\right)$  denoting a univalent acid radicle), but only the compounds of the last

type are capable of existing in aqueous solution, in which they are appreciably hydrolysed The solutions contain the colourless ion In . The ions In and In appear to be unstable and to undergo change as represented by the equations

$$3\ln^2 = \ln^2 + 2\ln^2$$
,  
 $3\ln^2 = 2\ln^2 + \ln^2$ 

Thus the lower halogen derivatives of indum are decomposed by water, metallic indium being deposited.

The tribables of indium appear to resemble the corresponding compounds of cadminin in their ability to form complex amons in solution, a resemblance which is not surprising since indium and eadmium occupy adjacent positions in the periodic table. The variation of the equivalent conductivity (A) tion (m, in gram-equivalents per litre) is shown for the four salts



Cube root of concentration in grm-eqs per litre with the cube root of the concentration (m. in gram-convergets per chloride and bromide solutions with those of cadmium bromide and nitrate

InCl<sub>3</sub>, InBr<sub>3</sub>, CdBr<sub>2</sub>, and Cd( $NO_3$ )<sub>2</sub> in fig. 17. The abnormally low values for  $\lambda$  in solutions of moderate concentration is shown in each case except that of cadmium intrate, and this salt does not form complex amons in solution. The low values for indium chloride are all the more remarkable since the data plotted refer to a temperature of 25°, the other data holding good for 18° C 4

Alloys.-Indium readily alloys with gallium,5 gold,4 lead,7 tin,8 and

Richards and Wilson, *loc cit* J. Hildebrand, J. Amer. Chem. Soc., 1913, 35, 501
 Heycock and Neville, Trans. Chem. Soc., 1889, 55, 666, 1892, 61, 911; see also \* Heycook and The Property of 
sodium,1 and easily forms an amalgam with mercury.2 When electrolytically deposited in a platinum dish it alloys with the platinum.3

#### INDIUM AND THE FLUORINE GROUP.

Indium Trifluoride, InF<sub>3</sub> -- Glistening, colourless, thombic pyramids of the trihydrate InF, 3H,O have been obtained by Thiel,3 and white needles of the hydrate InF<sub>3</sub> 9H<sub>2</sub>O have been prepared by Chabrié and Bouchonnet, by the action of dilute hydrofluoric acid on indium hydroxide. The hydrates are sparingly soluble in cold water, and when boiled with water are resolved into indium hydroxide and hydrofluorie acid.

The double salt endrum ammonium fluoride is very slightly soluble in

Indium monochloride, InCl, prepared by heating the dichloride with excess of indium and distilling the product in carbon dioxide, is a dark red solid which melts to a blood-red liquid. Its vapour density at 1100° to 1400° is about 5 3 (air - 1), corresponding to the simple formula " Water decomposes it into the trichloride and metallic indium

Indium dichloride, InCl., prepared by heating indium in a stream of dry hydrogen chloride, is a white, crystalline solid which melts to a yellow liquid. Its vapour density at 1167° to 1400° is about 6.5 (au. 1), corresponding to the simple formula? Water decomposes it into the trichloride and metallie indium.

Indium trichloride, InCl., is prepared by the action of excess of chloring upon indium, the lower chlorides, or a mixture of indium sesqui oxide and earbon, the product being purified by distillation in a stream of carbon It forms white, lustrous tablets, which do not volatilise appreciably at 410° Volatilisation occurs slowly at 600. The vapour density is 78, referred to an as unity, at 600 to 850°, corresponding to the molecular formula InCl<sub>2</sub>, but at higher temperatures dissociation occurs 8

The density of the solid chloride is 10.) The chloride is very deliquescent and dissolves readily in water, in which it is slightly hydrolysed, but the solution loses very little hydrogen chloride when evaporated at 100°.

The equivalent conductivities ( $\lambda$ ) of aqueous solutions of indium trichloride at 25° are, according to Thiel, as follows ( $\eta = {
m conc}$  in gram-equivalents per c c ; r = dilution, in litres per gram equivalent, see p. 155 and fig. 17) -

| 71         | 0.33 | 3 33 | 33 3 | 333   | 3333   |
|------------|------|------|------|-------|--------|
| $1000\eta$ | 3 () | 0.30 | 0.03 | 0.003 | 0.0003 |
| λ΄         | 10.2 | 30.5 | 50 6 | 101.0 | 2250   |

Indium trichloride forms double salts with the chlorides of the alkali metals, ammonium, and platinum.10 The salt KgInCl, 2H,O crystallises in the

Heycock and Neville, thid, 1889, 55, 666.
 Chabrić and Rengade, Compt. rend, 1901, 132, 472. Richards and Wilson, loc. ett.; J. Hildebrand, loc. cit.

Thiol, Zeitsch. unory Chem., 1904, 40, 280

Chabrie and Bouchonnet, Compt. rend, 1905, 140, 90. Huysse, Zottsch. anal Chem., 1900, 39, 9. Nilson and Pettersson, Trans Chem. Soc., 1888, 53, 814

Nilson and Petterson, loc. cit C. and V. Meyer, Ber, 1879, 12, 609, Nilson and Petterson, loc. cit.

R. E. Meyer Annalen, 1869, 150, 137; Nilson, Ber., 1876, 9, 1059, 1142.

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ditetragonal, bipyramidal class of the tetragonal system; and the ammonium, rubidium, and cessium salts of the type X,hrl, HO are isomorphous with one another, crystallising in the bipyrainidal class of the thombic system. The platinum salt has the composition 2InCl, 5PtCl, 36H,O. Indium trichloride also forms an addition product with animonia, and with pyridine forms the

Compound InCl<sub>3</sub> 3C<sub>5</sub>H,N, in p 253 <sup>2</sup>

Indium oxychloride, InOCl, obtained by the action of oxygen and chlorine upon heated indium dichloride, is a white, sparingly soluble powder.3

Indium monobromide, InB1, and indium dibromide, InB1, resemble the corresponding chlorides At 1.330° the vapour of the monobrounde has the normal density, but the dibrounde is considerably dissociated into the monobrounde and bromme 3

Indium tribromide, InBr, is prepared by the action of bromine vapour npon heated indum, and may be sublimed in a current of carbon dioxide resembles the trichloride in properties. Numerous physical properties of indium tribronide solutions have been incasured by Heydweller. The equivalent conductivities at 18° are as follows

| $1000_{\eta}$ $\lambda$ | 9 9701 | 5 0000 | 2 0032 | 0 9962 | 0 5003 | 0 2640 |
|-------------------------|--------|--------|--------|--------|--------|--------|
|                         | 0 1003 | 0 2000 | 0 1992 | 1 0038 | 1 9986 | 3 7875 |
|                         | 53 85  | 46 75  | 37 02  | 28 67  | 19 83  | 11 04  |

The double salts K  $\ln \mathrm{Br}_{\rm b}(2H_1O_1) \ln \mathrm{Br}_{\rm b}(H_1O_1) \ln \mathrm{Br}_{\rm b}(H_2O_1)$  and  $\mathrm{Cs_2InBr}_{\mathrm{c}}$  H O are isomorphous with the corresponding chlorine compounds

The following densities (at 20) and crystallographic constants are given by Wallace =

| Tetragonal | $\frac{\text{K}_4 \text{InCl}_6  2 \text{H}_2 \text{O}}{\text{K}_4 \text{InBr}_6  2 \text{H}_2 \text{O}}$                                                                                                                                 | a c        | I            | 0.81 | 7.3, |                              | density - | 2 483<br>3 140          |
|------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------|--------------|------|------|------------------------------|-----------|-------------------------|
|            | (NH <sub>1</sub> ) <sub>2</sub> InCl <sub>1</sub> H <sub>3</sub> O a<br>Rb <sub>2</sub> InCl <sub>2</sub> H <sub>3</sub> O<br>Cs <sub>2</sub> InCl <sub>2</sub> H <sub>3</sub> O<br>(NH <sub>1</sub> ) <sub>2</sub> InBi H <sub>3</sub> O | <i>h</i> , | 0.97         | (2.) | 1    | 1 4005 ,<br>1 4085<br>1 4033 | density - | $\frac{3089}{3350}$     |
|            | Rb,InBr, H <sub>2</sub> O<br>Cs <sub>2</sub> InBr, H <sub>2</sub> O                                                                                                                                                                       |            | 0 98<br>0 97 |      |      | 1 3951<br>1 4180             |           | 3 167<br>3 109<br>3 776 |

These results should be compared with those for the corresponding thallning salts (p. 181)

Indium oxybromide, InOBi, has been described a Indium mono-iodide, InI, is produced when an excess of indium is heated with iodine. Small quantities must be used, unless the reaction is carried out in an atmosphere of carbon dioxide - Indium tirrodide is reduced to the mono-iodide by repeated distillation in hydrogen - It is a brownish-red solid, which melts at  $351^{\circ}$  and boils at c 700', and may be distilled in carbon dioxide.

Indum mono include is not attacked by hot water, but dissolves in dilute

Wallace, Zedsch. Kryst. Mon., 1911, 49, 417
 Renz, Ber., 1901, 37, 2110., Dennis and Geer, ibid., 1901, 37, 961.
 Thiel, loc. cit

<sup>Heydwiller, Zeitsch, anorg Chem., 1911, 88, 103, see also p. 155 and fig. 17
Wallace, loc cit.
R. E. Meyer, loc cit., cf. Thuel, loc cit.</sup> 

acids with the evolution of hydrogen It is insoluble in alcohol, ether, and chloroform. The simultaneous action of air and water leads to the production of indum hydroxide and hydrodic acid. This action is slow with cold water, and the hydroxide is obtained in colloidal solution, but with hot water the hydroxide separates in a form easily filtered. 1

Indium di-iodide,  $\ln l_2$ , has not been isolated in the pure state, but thermal analysis shows that the compound exists. A mixture containing indium and iodine in the ratio In  $l_2$  melts at c 212, and the addition of either iodine or indium depresses the melting point. The failure to obtain the di-todde by heating the elements together in the correct ratio and distilling the product is doubtless due to dissociation of the molten di-iodide into the mono and triodides taking place to a very large extent, as the shape of the melting point curve indicates.2

Indium tri-iodide, InI3, is produced when indium is heated with an excess of todine in an atmosphere of carbon dioxide. It is a hygroscopic, yellow solid, melting at e 200° to a red liquid which can be slowly distilled in carbon dioxide at high temperatures. The tri-rodide is soluble in water, alcohol, ether, chloroform, and xylene

Indium perchlorate, In(ClO<sub>4</sub>), 811,0, prepared from indium and dilute perchloric acid, forms colourless, deliquescent crystals, which melt at 80° to a turbid liquid. It is easily soluble in water and alcohol.

Indium iodate, In(IO3), obtained from indium trichloride and potassium iodate, separates from its solution in hot dilute nitric acid as a white, crystalline powder, 1 part of which dissolves in 1500 of water at 18°, and in 150 of dilute intric acid (1.5) at 80°.4

#### INDIUM AND THE OXYGEN GROUP

Indium sesqui-oxide, In O3, obtained when indium is heated in air or oxygen, is prepared by the ignition of the hydroxide, carbonate, intrate, or sulphate Prepared at fairly low temperatures, it forms a yellow, amorphous powder which is readily soluble in acids, producing indium salts, but when produced at high temperatures, it forms rhombohedral crystals of the colour of chlorine, and is extremely resistant towards acids. The density is 7.18; the specific heat (0° to 100°), 0 0807.5 It is diamagnetic 5

Indium sesqui-oxide is reduced to indium when heated with sodium, carbon, or magnesium, or when heated in either hydrogen or ammonia, and by carefully regulating the reduction in hydrogen, Winkler claims to have obtained lower oxides, including a black monoxide InO

When heated above 850°, indium sesqui-oxide loses oxygen and becomes converted into the oxide In<sub>3</sub>O<sub>p</sub> which crystallises in regular octahedra isomorphous with Fe<sub>3</sub>O<sub>4</sub>.<sup>7</sup> The loss in weight associated with this change had previously been attributed to the volatility of the sesqui oxide b

<sup>&</sup>lt;sup>1</sup> Thiel and Koelsch, Zeitsch. anong Chem., 1910, 66, 288.

<sup>2</sup> Thiel and Koelsch, loc at large Thiel and Koelsch, loc at large R. E. Mayer loc. cit., Thiel, loc atl., Thiel and Koelsch, loc. cit.

<sup>4</sup> Mathers and Schluederberg, J. Amer. Chem. Soc., 1908, 30, 211

<sup>5</sup> Nilson and Petersson, Compt. rend., 1880, 91, 232; Ber., 1880, 13, 1459

<sup>6</sup> Winklet, J. prakt. Chem., 1865, 94, 1, 95, 414, 1867, 102, 273, and Ber., 1890, 23, 772; Dennis and Geer, J. Amer. Chem. Soc., 1904, 26, 437; Ber., 1904, 37, 93.

<sup>7</sup> Thiel and Koelsch, loc cit., cf. Rend, Trans. Chem. Soc., 1894, 65, 313.

<sup>8</sup> Renz., Ber., 1908, 36, 1847; 1904, 37, 2110; Meyer, Zeitsch. anorg. Chem., 1905, 47, 281; Thiel, ibid., 1906, 48, 201.

Neither the oxide In O4 nor the (metastable) In O3 fuses in the oxyhydrogen flame

From the data supplied by Intte 1 the following results may be deduced -

$$\begin{array}{ll} 2[\ln] + 3(0) &= [\ln_2 O_3] + 239 \ 8 \ \mathrm{Cals}, \\ [\ln_2 O_3] + 3 \mathrm{H}_2 \mathrm{S} O_4 \ \mathrm{aq} &= \ln_2 (\mathrm{S} O_4), \ \mathrm{aq} \ + 72 \ 78 \ \mathrm{Cals}. \end{array}$$

Indium hydroxide, In(OII),—Indium is slowly converted by air and moisture into indium hydroxide. The hydroxide may be prepared by adding ammonia to an aqueous solution of an indinin salt and washing the precipitate. Other precipitants may be used instead of ammonia, eg. potassium nitrite, hydroxylamine, and the simple aliphatic primary and secondary amnes. The air-dired precipitate has the composition 21n(OH), 3H<sub>2</sub>O, and loses 3H<sub>2</sub>O at 100° 3

Freshly precipitated indimin hydroxide exhibits a marked tendency to pass into the colloidal state in the absence of electrolytes. It is slightly soluble in concentrated ammonium hydroxide, and dissolves readily in alkali hydroxides From the latter solutions it is reprecipitated on boring or standing. When heated to redness, it leaves indinin sesquioxide. With ddinte acids it reacts to produce indium salts and water, but it possesses slight acidic properties, a magnesium indate being known. This substance, of the formula Mg ln<sub>2</sub>O<sub>1</sub> 3H<sub>2</sub>O<sub>2</sub> is obtained as a white precipitate when aqueous solutions of magnesium and indium chlorides are mixed and boiled by

Indium monosulphide, In S, may be prepared by heating the trisulphide in a stream of hydrogen. It is volatile at a red heat and forms flat, transparent, yellowish-brown crystals b

Indium disulphide, InS, is prepared by heating indium in a stream of hydrogen sulphide, and forms a soft, reddish brown solid 6

Indium sesquisulphide or trisulphide, In S, -This compound was prepared by Winkler as a brown solid by heating either indium or its sesqui-oxide with sulphur. By igniting a mixture of sodium carbonate, indium oxide and sulphur, Winkler obtained the sesquisilphide of indium in shining, yellow crystals. Thiel has prepared the sulphide in searlet-red shining crystals by igniting indium sesqui oxide in a current of hydrogen sulphide. The sulphide is not volatile at a red heat, but when heated in air it is decomposed with the ultimate formation of indium sesqui-oxide?

When hydrogen sulphide is passed into an aqueous solution of an indium salt, indium sesquisulphide is thrown down as a yellow precipitate, resembling cadmium sulphide in appearance. The precipitation is prevented by the presence of immeral acids except when the acidity of the solution is very slight, but acctic acid does not inhibit the precipitation The yellow sulphide is soluble in concentrated mineral acids

Indium sesquisulphide is obtained in colloidal solution by passing hydrogen sulphide into water in which indium hydroxide is suspended. Excess of

Intic, Compt. rend., 1871, 72, 858., 73, 108
 Ruz, Ber., 1901, 34, 2763, 1903, 36, 2751; 1904, 37, 2110, Paper., Annalen, 1871, 158, 372; Dennis and Geet, Ber., 1904, 37, 901. J. Amer. Chem., Noc., 1904, 26, 437.
 Winkler, J. prakt. Chem., 1867, 102, 273, Cainciley and Walker, Trans. Chem. Soc., 1888, 53, 74, 88
 Renr, Ber., 1901, 34, 2763
 Thiel, Zertoh. anory. Chem., 1904, 40, 280, Thiel and Koelsch, ibid., 1910, 66, 288.
 Thiel and Koelsch, loc. cit.
 Winkler, J. prakt. Chem., 1867, 102, 273; Thiel, Zeitsch. anory. Chem., 1904, 40, 280.

hydrogen sulphide may be removed by boiling, but the addition of an electrolyte leads to the precipitation of the yellow sniphide

Indium sesquisulphide forms either double or complex salts with alkali sulphides Indiam potassium sulphide, K2SAn2S3 or K2(In2S4), is obtained as insoluble, hyacinth red, quadratic tables when indium sesqui-oxide (1 pt ), potassium carbonate (6 pts.), and sulphur (6 pts.) are fused together using sodium carbonate, the product obtained is soluble in water, but a white precipitate of indian sodium sulphide, Na<sub>2</sub>(In<sub>2</sub>S<sub>4</sub>) 2H<sub>2</sub>O, slowly separates out <sup>2</sup>

Potassium sulphide gives a white precipitate, probably of the composition  $\mathbf{K}_{2}(\operatorname{In}_{2}S_{1})/(\operatorname{II}_{2}O_{1})$  when added to the solution of an indium salt. In excess of potassium sulphide the precipitate dissolves. The addition of immeral acid to this solution leads to the precipitation of yellow indium sulphide, which dissolves in excess of the acid. Potassium hydrosulphide also produces a white precipitate with an indium salt, but it is insoluble in excess of the precipitant. An analogous white precipitate is produced by ammonium sulphide, it dissolves in excess of the hot, but not of the cold precipitant?

Basic indium sulphite, In<sub>2</sub>(SO<sub>3</sub>)<sub>3</sub> 2In(OH)<sub>2</sub> 5H<sub>3</sub>O<sub>3</sub> is obtained as an insoluble, white precipitate when a solution of an indium salt is boiled with sodium or anunonium sulplite. It dissolves in cold aqueous sulphirous acid, but separates out again as a crystalline powder when the solution is boiled It loses 311,0 at 100°, and another 511,0 at 260°. At 280 sulphur dioxide begins to be evolved, and when further heated a residue of indium sesqui-oxide is ultimately obtained 1

Indium sulphate, In (801), may be prepared by dissolving indium, indium sesqui oxide, or indium hydroxide in sulphinic acid, and carefully heating to expel water and excess of sulphuric acid. It is a white solid, very hygroscopic and extremely soluble in water, from which it is exceedingly difficult to crystallise out a hydrated sulphate. An ennea hydrate,  $In_{g}(SO_{4})_{g}9H_{2}O_{5}$  is said to exist, and an acid salt of the composition  $In_2(SO_1)_1 Il_2SO_1 8Il_2O$  is described as separating from an acid solution of the sulphate when evaporated over concentrated sulphuric acid."

The anhydrous sulphate has a density of 3.11 and a specific heat (between 0° and 100') of 0 129 6

Various double sulphates have been prepared by mixing indium sulphate solutions with solutions of other sulphates. The ammonium-, rubidium-, and casium-indium alums, MaSO<sub>4</sub> In<sub>2</sub>(SO<sub>4</sub>), 24H<sub>2</sub>O (Ma-NH<sub>2</sub>, Rb, or Cs), can be thus prepared, the first having been discovered by Roessler and the others by Soret Lithium, sodium, and potassum alims are not known, but sodium indium sulphate, Na,SO, In,(SO,), SH,O, and the corresponding potassium, ammonium, and thallium salts have been prepared

The three alims separate out in regular octahedra or condinations of the octahedron and cube. Then aqueous solutions have an acid reaction, and when heated become turbid owing to the separation of basic sulphates

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Winssinger, Bull. Soc. chim., 1888, [n.], 49, 152, Linder and Pieton, Trans. Chem.
 Soc., 1892, 61, 134
 Schmeider, J. piakt. Chem., 1874, [n.], 9, 209.
 Rechand Richter, J. piakt. Chem., 1803, 90, 172, 1864, 92, 480
 Winkler, thirl., 1867, 102, 273, R. E. Meyer, Annalen, 1869, 150, 137
 C. E. Bayer, Annalen, 1871, 188, 372; Ann. Chem. Phys., 1872, [iv.], 23, 50, Thiel, Zettsch. annag. Chem., 1904, 40, 280.
 R. E. Meyer, Annalen, 1869, 150, 137
 Wilson and Patterson, Ext. 1880, 12, 1489, Count, 2nd. 1880, 01, 232

<sup>\*</sup> Nilson and Pettersson, Ber., 1880, 13, 1459; Compt. rend., 1880, 91, 232.

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or, in the case of the cassium salt, indium hydroxide. The melting points and solubilities of the alums in water (in grains of crystalline salt per 100 grams of water) are as follows .- 1

|            |        |      |    |      |      |       | Solub | ilitics. |        |
|------------|--------|------|----|------|------|-------|-------|----------|--------|
| Temperatur | :e, *C |      |    |      |      | 15°   | 16 5° | 25°      | 30°    |
| Ammonium   | ındıum | alun | n, | m.p. | 36°, |       | c 200 |          | c. 400 |
| Rubidium   | **     |      |    |      |      | 41 28 |       |          |        |
| Cæsium     | ,,     | ,,   | ,  | mр   | ,    |       | 3 04  | c. 117   | •      |

In the case of the ammonium alum, the "melting-point" is known to denote the transition-point, above which the alum breaks up into the octahydrate and water (Roessler) -

$$(NH_4)_2SO_4 In_2(SO_4)_3 24H_2O \longrightarrow (NH_4)_2SO_4 In_2(SO_4)_3.8H_2O + 16H_2O$$

The densities and refractive indices of the alums for the D-line at the ordinary temperature are given by Soret 2 as follows :-

|         | NH.    | Rb     | Cs     |
|---------|--------|--------|--------|
| density | 2011   | 2 065  | 2.211  |
| μ       | 1 4664 | 1 4638 | 1 4652 |

Indium sesquiselenide, IngSeg, is obtained by heating indium with selemum and repeatedly fusing the product with selemum. A brown precipitate of the same composition is produced by passing hydrogen selenide into indium a etate solution. The sesquiselenide is decomposed by dilute hydrochloric acid with the evolution of hydrogen selende, and is eventually converted by mtric acid into indium selemite. When heated in hydrogen, hydrogen selende and a dark sublunate (probably a lower selende) are produced

Indium selenite,  $\ln_2(\text{SeO}_3)_3$  6H<sub>2</sub>O. -When sodium selenite is added to a solution of an aluminum salt, a basic sciente is precipitated, which is converted into the normal selemite when heated with aqueous selemous acid at 60. The normal sciente is a white, crystalline powder, practically insoluble in water. Two acid selenites have been also described

Indium selenate, In<sub>2</sub>(SeO<sub>4</sub>)<sub>3</sub> 10H<sub>2</sub>O<sub>5</sub> is prepared by dissolving indium hydrovide in aqueous science and and crystallising the solution. The crystals are hygroscopic and easily soluble in water. When mixed with the requisite amount of aqueous cosmin selenate, cæsium indium selenate, lu2(SO4), Cs2SO4 24H2O, crystallises out in efflorescent crystals, which are described as tetragonal bipyramids and not regular octahedia as would be anticipated 5

Indium telluride, InTe, may be prepared by fusing together indium and tellurium in an atmosphere of nitrogen. It forms a readily fusible solid, which is insoluble in hydrochloric acid but attacked by intic acid. A brown indium telluride is precipitated when hydrogen telluride is passed into a solution of indium acetate

Roessler, J. prakt Chem., 1873, [11], 7, 14, Chabné and Rengade, Compt. rend., 1900, 131, 1300, 1901, 132, 472; Bull. Soc. thin., 1901, [10], 25, 566, Locke, Amer. Chem. J., 1901, 26, 166.

<sup>01, 20, 180.

&</sup>lt;sup>3</sup> Soret, Arch. Scs. phys. nat., 1885, [m ], 13, 5, 1888, [m.], 20, 520.

<sup>5</sup> Thel and Koelsch, Zedsch. ann. J. Chem., 1910, 66, 288

<sup>4</sup> Nilson, Nova Acta Soc. Upsatt, 1875, [m ], 9, No. 7, Ball. Soc. chem., 1875, [n ], 23, 191

Mathers and Schluederberg, J. Amer Chem. Soi., 1908, 30, 211.
 Thiel and Koelsch, loc. cut.

Indium chromate does not appear to have been analysed Indium molybdate,  $In_2(MoO_4)_3$   $2H_2O$ , indium tungstate, and indium uranate are mentioned by  $Ren^{2/3}$ 

Indium silicotungstates. Indium silicotungstate is very soluble in water, and two hydrates may be crystallised out from the solution. The first, of the composition  $2\ln \rho_0^2 3(\sin \rho_1) 2WO_3 93H_2O$ , forms regular octahedia, while the second, with  $63H_2O$ , is trichine  $(a,b,c=0.6025-1-0.7864+a=85^\circ-10',\beta=106^\circ-3',\gamma=90'-12')$  The acid salt,  $\ln \rho_0$ ,  $H_2O_2(\sin \rho_1) 2WO_3 40H_2O$ , forms monochine crystals  $(a-b-c=0.9552:1.0^\circ6544+\beta=96-32')^2$ 

#### INDIUM AND THE NITROGEN GROUP.

Indium nitrate,  $2\ln(NO_3)$ ,  $9H_3O$  — Indum dissolves slowly in dilute, more rapidly in concentrated intric acid, a solution of the intrate being produced. In the presence of intric acid, indium intrate readily crystallises from the solution, but from a neutral solution it is very difficult to crystallise the salt. Indum nitrate forms colomiess, deliquescent needles, it loses one third of its water at 100 and all of it over concentrated sulphuric acid The intrate is easily decomposed by heat, and its aqueous solution becomes turbid, owing to the separation of basic salt, when heated (Winkler) A double untrate is formed with ammonium intrate 3

Indium phosphide, InP, is left as a black, crystalline residue when indium mono jodde is melted with white phosphorus and then heated to 400' Metallic indium is only superficially attacked by phosphorus 1

Indium phosphate. - The precipitate obtained by adding sodium phosphate to an indum salt has not yet been examined in detail

Indium vanadate. - Indium salts are precipitated by ammonium metavanadate 1

## INDIUM AND THE CARBON GROUP.

Indium carbonate (!) has not yet been analysed

Indium cyanide also does not seem to have been analysed.

The platinocyanide, In [Pt(CN)<sub>1</sub>], 211<sub>2</sub>O, forms white, hygroscopic, soluble leaflets

Indium oxalate, ln<sub>2</sub>(C<sub>2</sub>O<sub>4</sub>)<sub>3</sub> 6H<sub>2</sub>O<sub>5</sub> is insoluble in water and ammonia <sup>5</sup> Other organic salts of indium have been prepared by R. E. Meyer

Indium acetylacetonate, [(CII, CO),CII], in, is prepared by heating indum hydroxide for eight hours with a boiling alcoholic solution of acetylacetone. It erystallises from alcohol in flat, six-sided prisms and inelts at 183°, the crystals are rhombic (bipyrainidal) and isomorphous with scandium and ferric acetylacetonates. At 260° to 280° it partly sublines and partly decomposes. Its molecular weight in ethylene dibromide solution is in accordance with the above simple formula,8

Renz, Ber., 1901, 34, 2763
 Wyrouboil, Bull. Soc. trans. Min., 1907, 30, 277
 Dennis and Geer, J. Amer. Chem. Soc., 1904, 26, 437.
 Thell and Koelsch, loc. cit.
 W. M. J. Chem. Chem. 1807, 200, 273, Bettern.

Winkler, J. prakt. Chem., 1867, 102, 273, Bottger, thid., 1866, 98, 26. Huysse,
 Zeitsch and Chem., 19, 0, 39, 9.
 R. E. Meyer, analen, 1869, 150, 137
 Jaeger, Proc. R. Akad. Wittensch. Amsterdam, 1914, 16, 1095, Revitar chim., 1914,

<sup>33, 342</sup> Chabné and Rengado, Compt. rend., 1900, 131, 1300, 1901, 132, 472.

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#### DETECTION AND ESTIMATION OF INDIUM.

Indium salts colour the flame bluish-violet, and spectroscopic examination of the flame reveals the two dominant lines \$4511.5 and \$4102.0 To detect indium in a immeral, it may be attacked with aqua regia, the acid neutralised, and, after adding excess of sodium acetate, the solution saturated with hydrogen sulphide. The precipitated sulphides may then be treated with hydrochloric acid, and a spectroscopic test carried out 1

Heated in the reducing flame on charcoal with fusion nuxture, indium compounds give a metallic bead and a yellow incrustation of oxide

Indium hydroxide, freshly precipitated, is only appreciably soluble in an excess of concentrated ammonium hydroxide. It dissolves in sodium or potassium hydrovide, but is reprecipitated on boiling the solution or on standing Indium is completely precipitated as the hydroxide by barmin Indium carbonate (') is soluble in cold ammonium carbonate solution, but is reprecipitated on boiling the solution. Indiam salts are precipitated by potassium cyanide, ferrocyanide, or chromate, but not by the dichromate. The precipitated cyanide is soluble in excess of the precipitant, but on standing or by boiling, indinin hydroxide separates from the solution. Indiana ferrocyanide is much more soluble in hydrochloric acid than is the corresponding gallium salt

In neutral solutions, or solutions acidited with acetic acid, indium is completely precipitated as the yellow solphide. In solutions containing mineral acid (except in very low concentration) the precipitation does not occur Alkalı sulphides produce more complex precipitates

Very small quantities of indium, may be detected interochemically by the formation of inbiding indum chloride -

Indium is precipitated quantitatively from its salts as the hydroxide by boiling with a slight excess of ammonia. The precipitate is dired at 850° and weighed as the sesqui-oxide 8. In place of animonia as precipitant, directly lanime has been suggested, and also methylamine and ethylamine or their hydrochlorades, while Stock's method of precipitation (see p. 100) may be adopted

Indium is difficult to separate from iron. The best method appears to be to deposit the indium electrolytically from its sulphate solution, dissolve the deposit, which will still retain a little non, in intric acid, nontralise with ammonia, add an equal volume of 50 per cent, acetic acid, and procepitate the from by the addition of a solution of introso- $\beta$  naphthol in acetic acid

For the electrolytic deposition of indium, a sulphate solution containing a little free acid may be employed. Third uses a silvered platinum bowl as cathode, rapid deposition may be obtained with a merciny cathode and rotating anode. Good deposits on platinum cathodes are also obtained from warm cyanide or tartrate solutions, employing a rotating anode "

Hoppe-Scyler, Annalos, 1866–140, 247.
 Kley, Chem. Zett., 1901, 25, 563., of Haysse, Zeitsche anal. Chem., 1900, 39, 9.
 Ibiel and Koelsch, Zeitsche anorg. Chem., 1910, 65, 288.
 Henz, Berr., 1901, 34, 2763., 1203, 36, 2751, 4494.
 Mathers, J. Amer. Chem. Soc., 1908, 30, 209.
 kollock and Smith, ibid., 1910, 32, 1248.

### CHAPTER VIII.

#### THALLIUM.

Symbol, TI Atomic weight, 204 0 (0 = 16).

Occurrence. - Thallium is one of the narer elements, and only two or three inmerals are known in which thallium is an essential constituent. Crookeste, an apparently amorphous mineral discovered by Nordenskiold in a copper mine at Skrikerum (Sweden), has the composition Tl 17-25, Se 33-28, Cu 45-76, and Ag 3.71 per cent <sup>1</sup> Lorandite, TlAs><sub>2</sub>, has been found as red, translucent, monochine crystals  $(a \cdot b : c = 1.3291 - 1.0780 : \beta = 52^{\circ} - 27')$  m Macedonia, associated with realgar 2 Vrbaite, TlAs, SbS, also occurs in Macedonia, associated with realgar and orpiment - It forms orthorhombic crystals  $(a, b \ c = 0.5659 \ 1 \ 0.4836)$  which are deep red and translucent when small, and give a red streak.3

In small quantities thallium occurs widely distributed in nature in rocks and minerals, sometimes in association with the alkali metals, at other times with non, zinc, lead, etc 4 Thus it occurs in (a) lithia mica, (b) sylvine and carnallite, (c) alumte, (d) zinc blende, (e) pyrites, hiematite, and marcasite, (f) braumite, (g) sphalomite, etc 5 Thallium is also found in certain mineral waters, eg at Nauheim, and is widely diffused in the vegetable kingdom

Owing to the occurrence of traces of thallium in various minerals, it is found in small amounts in a number of commercial products, eg zinc, cadmium, platmium, bismuth, tellurium, etc  $^7$ —It also occurs in the chamber mud and flue dust from vitriol factories where thalliferous pyrites is burnt

<sup>1</sup> Nordenskield, K. Stenska Vet .- Akad Handl., 1866, No. 10; Annalen, 1868. 145,

Nordensktold, K. Nienska Vet.-Akad Handl., 1866, No. 10; Annalen, 1868. 145, 127
 Kienner, Zeitsch Kryst Min., 1897, 27, 98., Goldschnudt, ibid., 1898, 30, 272
 Jerek, ibid., 1912, 51, 361, Kichlik, ibid., 1912, 51, 379
 Schramm, Annalen. 1883, 219, 374, Hartley and Ramage, Trans Chem. Na., 1897, 71, 533; Vernadski, Hull. Acad Net. Petrograd, 1909, p. 821.
 (a) Schiotten, Sitzungsber K. Akad. Wiss. Wien, 1863, 48, 734, 1864, 50, 208; J. prakt Chem., 1864, 91, 45, 93, 275, (b) Hammerbacher, Annalen, 1875, 176, 82; (c) Cosa, Gazetta, 1878, 8, 235; Ber., 1878, 11, 811, (d) Bunsen, Annalen, 1865, 133, 108; Ulbam, Compl. rend., 1909, 149, 602, (c) Cuokes, Lamy, Kuhlmann, vode infra, Playfair, Chem. News, 1879, 39, 245, Antipoff, J. Russ. Phys. Chem. So., 1896, 28, i. 384; Inglestrom, Zeitsch Kryst Min., 1895, 25, 94; (f) Bischoff, Annalen, 1864, 129, 375; (g) von Kobell, J. prakt. Chem., 1871, [n], 3, 176.
 Bottger, Annalen, 1861, 127, 368, 128, 240.
 Kosmann, Chem. Zeit., 1886, 10, 762; Pinpson, Compt. rend., 1871, 78, 563, Watten, Chem. News, 1887, 55, 241; Horepath, ibid., 1863, 7, 77, Phurm. J., 1863, 4, 302; Crookes, Chem. News, 1863, 7, 109; Werther, J. prakt. Chem., 1803, 88, 180, Roepper, Amer. J. Sci., 1863, [n.], 35, 420; Streng, Dingl. poly. J., 1865, 177, 329, Zunmermann, Annalen, 1886, 232, 278.

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History.—Thallmin was discovered independently by Crookes in England and Lamy in France. Each of these scientists observed a new green line in the spectrum of some material he was examining, attributed it to a new element, and succeeded in isolating it. Crookes, who in March 1861 was the first to make the discovery, was engaged in extracting selemum from a deposit obtained from a sulphuric acid factory at Tilkerode in the Haiz. At first he suspected that the new element was a metalloid belonging to the sulphur group, and called it thallium in allusion to the green colour of its spectrum line (Latin, thallus, a budding twig)—His early work on thallum was greatly hampered by lack of material, but he eventually discovered the metallic nature of thallium, and in May 1862 was able to exhibit a few grains of the metallic element in powder form 1. Lamy made his discovery in April 1862, when examining the lead chamber deposits from the sulphuric acid factory of M F. Kuhlmann at Loos, where Belgian pyrites were burnt. Being more fortunate than Crookes in having considerable material at his disposal, Lumy was able very quickly to establish the metallic nature of thallium and to exhibit a himp of the metal in May 1862, and before the end of the year he was able to isolate several hundred grams of thallum and give a tarrly complete account of the physical and chemical properties of the element He showed that thallium forms more than one series of compounds, and that one series closely resembles the corresponding series of compounds of the alkali metals. Further, he found that in this series of compounds the equivalent of thallium is about 204, and (with the assistance of Regnault) showed by the application of Dulong and Petit's Law that this number also represents the atomic weight of thallinn. During the same year Kuhlmann prepared a number of organic thallium salts, and de la Provostaye indicated the isomorphism of several of them with the corresponding salts of potassium. During the next few years a considerable amount of work on thallium was published, notably by Crookes, Lamy, Desclotzeaux, Kuhlmann, Weither, Bottger, Nicklés, Willin, and Carstanjeu

At the time when thallium was discovered, the Periodic Classification had not been formulated, and it proved a difficult matter to place thallium in the classification generally accepted at that period for the metals. A number of thallous compounds were observed to resemble closely the corresponding compounds of lead in their physical properties, and even more remarkable was the resemblance between the actual elements themselves, lead and thallium. Other thallous salts, however, were observed to be isomorphous with, and to rescuble quite closely, the corresponding salts of potassium. Moreover,

Chookes, Chem. News, 1861, 3, 193, 393, Phil. May, 1861, [iv], 21, 391.
 Lamy, Soc. Imperante des Sennes de Lette, May 16, June 20, Aug. 1, Sept. 5, Nov. 7, 1862, Compd. rend., 1862, 54, 1255, 55, 836, Ann. Chem. Phys., 1863, [in.], 67, 385; Regnault, Compt. rend., 1862, 55, 887, Ann. Chem. Phys., 1863, [in.], 67, 427, Kuhlmann, Compt. rend., 1862, 56, 607, 1863, 56, 171, Ann. Chem. Phys., 1863, [in.], 67, 428, 431; de la Provostaye, Compt. end., 1862, 55, 610
 Crookes, Chem. News, 1862, 6, 1, 1863, 7, 109, 133, 115, 194, 218, 290, 8, 150, 195, 219, 231, 243, 255, 279; 1864, 9, 1, 37, 51, 1867, 15, 294, Phil. May., 1803, 26, 55, Proc. Roy. Soc., 1862-63, 12, 150, Crookes and Church, Chem. News, 1863, 8, 1, Lamy, Ann. Chim. Phys., 1864, [in.], 3, 33, 1865; [iv.], 5, 410, Lamy, and Descloteaux, ibid., 1869, [iv.], 17, 310, Kuhlmann, Bull. Soc., them., 1864, [in.], 1, 330, 3, 57, Werther, J. prakt. Chem., 1863, 89, 189, 1864, 91, 385; 92, 128, 351; 93, 393, Bottger, Annaten, 1863, 126, 266; 127, 368, 128, 210, 218, Nikles, Compt. rend., 1851, 58, 192, 537, J. Thorm Chim., 1866, [iv.], 4, 127; Willin, Ann. Chim. Phys., 1865, [iv.], 5, 5, Carstanjen, J. prakt. Chem., 1867, 101, 55, 102, 65, 129.

the spectrum of thallium was extremely simple and very easy to observe, resembling in this respect the spectra of sodium, potassium, and the then newly discovered elements rubidium and cosmim—It is therefore not surprising that Dumas should have reported to the French Academy that "le thallium offre une rémnou de propriétés contradictores qui autoriserait à l'appelei le métal paradoxal, l'orinthorynque des métaux " !

To classify thallium with the alkali metals is clearly unsatisfactory, since the element itself is decidedly different in properties from sodnim, potassium, ete, and, moreover, it forms a series of salts in which it is tervalent, to classify it with lead is worse, since the only analogies between lead and thallium salts are physical analogies between salts of univalent thallium and bivalent lead With the publication of the Periodic Classification, Mendeléeff pointed out that thallium should really be classified with aluminium and indiam, and made out a very good case for this contention? Recalling the fact that when the elements are arranged in the increasing order of their atomic weights, aluminium comes between magnesium in Group 11 and silicon in Group IV, while thallium comes between mercury in Group II and lead in Group  $\dot{IV}$  , the case may be stated almost exactly as it was given by Mendeléefl Only the highest oxide of mercury shows any analogy with magnesium oxide; only the highest oxide of lead shows any analogy with silica. In the same way, only the highest oxide of thalling shows any analogy with alumina Magnesium and mercuric oxides are basic oxides and give rise to salts of the type MX,, aluminum and thallic oxides are more feeble bases and give rise to salts of the type MX3, silicic and plumbic oxides are feeble acidic oxides Thallium gives, independently of thallie oxide, a lower oxide which is a powerful base, aluminium torms no lower basic oxide. Strictly analogous is the existence of basic exides of mercury and lead lower than mercuric and plumbic oxides, and the non-existence of lower oxides of magnesium and silicon The higher oxides of the formulae HgO, TLO, PhO, and Bi,O, are peroxides relatively to the lower oxides of the formula Hg<sub>2</sub>O, Tl<sub>2</sub>O, PbO, and Bl<sub>2</sub>O<sub>3</sub>, and give off oxygen when heated The higher chloride of mercury is stable, that of thallium mustable, the tetrachloride of lead very mustable, and pentachloride of bismuth non-existent. Bismuth trichloride is decomposed by water, and lead (di)chloride when heated in steam, but thallous chloride is stable. Thallum is less volatile than mercury but more so than lead. The similarity between thallous and alkali salts is no more remarkable, than that between the plumbons salts and the salts of the alkaline-earth metals, or that between bismith salts and the salts of certain elements of Group III, (see p. 234). There is, then, no more difficulty in placing thallium in Group III than there is in placing mercury in Group II, lead in Group IV, or bismuth in Group V.

Preparation .- The best source of thalhum is thalliferous non pyrites All deposits of pyrites do not contain thallium in appreciable quantity mentions the Belgian pyrites from Theux, Namur, and Philippeville and cortain Spanish pyrites as being particularly rich in thalloim

Methods for extracting thallinm from thalliferous pyrites have been described, but are rarely, if ever, used. Thallium is best obtained from the

<sup>1</sup> Dumas, Pelouze, and Deville, Report to the French Academy, Compt. rend., 1862, 55,

<sup>866;</sup> Ann. Chim Phus, 1863, [in ], 67, 418

Mendeleeft, Amalin Suppl., 1872, 8, 133, or Chem. News, 1879, 40, 231, etc., 1880, 41, 2, etc.; cf Otto, J. prakt. Chem., 1867, 102, 185, Eidmann, dud., 1863, 89, 381; Rammelsberg, Ber., 1870, 3, 276

flue dust or chamber deposits from sulphune acid works in which thalliferous pyrites is birint.

Flue dust, which contains thallinin, lead, arsenic, selenium, etc., is re-peatedly boiled with water slightly acidified with sulphuric acid. The filtered solution is concentrated and the thallium precipitated by introducing metallic zinc into the solution. Thallium is thereby precipitated in the torm of needles or glittering plates. Alternative procedures are to precipitate the thalloun as thallous chloride by adding a soluble chloride, or as thallous sulphide by adding sodium carbonate, filtering from precipitated imporities, adding potassium cyanide, and then saturating with hydrogen sulphide

Lead chamber deposit 2 is neutralised with lead oxide or lime and extracted with hot water. The solution is filtered and concentrated, and the thalloun precipitated as metal, chloride, or sulphide as described above

The zine sulphate mother liquors obtained from a white vitrol factory at Goslii in the Haiz were found by Bunsen to yield 0.05 per cent of their weight of thallous chloride. The thallium in such a solution may be precipitated by means of zinc, the deposit, containing a little cadmium and copper, treated with dibite sulphuric acid to dissolve the thalloun and cadminin, and the thalloun precipitated as thallous todide from the filtered solution

The crude metallic thallour is purified by converting it into thallous sulphate and treating the slightly acid solution with hydrogen sulphide to channate traces of mercury, silver, arsenic, antimony, and bismuth. The filtered solution is treated with ammonia to remove traces of non and aluminum, and the filtrate concentrated until thallons sulphate crystallises out Crude thallous chloride is slowly added to hot, concentrated sulphurio acid and the mass heated until all hydrochloric acid is expelled solution is, as in the preceding case, purified by successive hydrogen sulphido and annuous treatments, and thallous sulphate then crystallised from the solution. A smalar method of treatment may be applied to crude thallous sulphide. From the pure thallons sulphite, thallium is most conveniently obtained in a state of purity by electrolysis, as described by Foerster 5. pure thallous iodide (chloride or brounde), thallium may be prepared by fusion with potassium evanide and sodium carbonate (Werther). Thallium may be

also prepared by heating thallous oval ite in a covered criticible (Willin)

Properties. Thalloun is a blinish white metal. It is so soft that it can be seratched with the finger nail and cut with a knife. When freshly cut it exhibits a bright metallic histre. The metal is extremely malleable, but has little tenacity. Its structure is crystalline, and when a bar of thalloum is bent it enuts a sound resembling the "cry of tin". Thallium exists in two

Cookes, Chem. No.s, 1863, 8, 159. Bottger, Annalen, 1863, 126, 266. 128, 248;
 Carstanjen, Aid., 1867, 101, 55. 102, 65, 129. Schaffner, Dand. poly. J., 1872, 205, 55;
 Schangsher K. Akad. Wiss. Wien. 1871, 63, 176. Gunning, Bull. Soc. chim., 1868. [n.],
 10, 359. Aich. Noerland, 18-8, 3, 86. Nuclei, Aich. Pharm., 1875. [m.], 7, 385. Dangl.
 pala. J., 1876. 219, 262. Stellar, Chem. Soc. Ibd., 1874, 27, 873. Kituse, Dangl. polit. J.,
 1875, 217, 323. Weither, J. probl. Chem., 1804, 91, 385; 92, 128, 361; Wohler, Annalen,
 1867, 142, 263., 1872, 164, 74. Forester, Zeitsch. aneog. Chem., 1897, 15, 71.
 Luny, Ann. Chem. Phys., 1863, [m.], 67, 385.
 Bunsen. Annalen, 1865, 133, 108. Schumann, ibid., 1888, 249, 340. Bartlett, J.
 Soc. Chem. Lud., 1865, 133, 108. Schumann, ibid., 1888, 249, 340. Bartlett, J.
 Soc. Chem. Lud., 1865, 134, 186.
 On. the preparation of pure thallum see Crookes, Phil. Trans., 1873, 163, 277;
 Chem. News, 1874, 29, 14, etc.
 Foeister, Zeilsch. anorg. Chem., 1897, 15, 71.

modifications which are enantiotropic; the transition-point is 226° C, and is lowered 6° for an increase of pressure of 3000 kilos per sq cm. Accordingly, as the temperature is raised through 226° a contraction in volume occurs, amounting to 0 000011 cc per grain. The density of thallium is 11.85 at 20° C, 2 the coefficient of expansion (0° to 100°) is 0.00003135,3 and the compressibility coefficient is  $2.33 \times 10^{-6}$  per atmosphere.<sup>4</sup>

Thallium inelts at  $302^{\circ} \pm 1^{\circ}$  C.,<sup>5</sup> the volume increasing by 3.1 per cent.

during the change. The latent heat of fusion is 7.2 cals per gram 7 In vacuo, thallium commences to volatilise at 174°, and boils at 818° under a pressure of 15 cm of its own vapour 8 It is said to boil at 1515° under normal pressure " von Wartenberg, however, gives the following values for the vapour pressure of thallium.

```
Temperature, *('.
                          . 631°
                                      783°
   970° C
Vapour pressure .
                          0.056
                                      1 329
   2131 mm of Hg
```

and estimates the boiling-point to be 1306° 10. The mean specific heat of thallium between 20° and 100° is given as 0.0325, 0.0336, and 0.0326 by Lamy, Regnault, and Schmitz respectively, between -188' and +20' the mean value is given as 0 0300 by Schmitz and 0 0296 by Richards and Jackson <sup>11</sup> The specific heat is 0 0277 at  $-177.3^{\circ}$  and 0 0143 at  $-250 \cdot 1^{\circ}$  C <sup>12</sup>

The electrical conductivity (a) of thallium (in recipiocal ohms per cm. cube) at various temperatures is as follows — 13

Temperature, °C. 
$$-180^{\circ}$$
  $-80^{\circ}$   $-10^{\circ}$   $0^{\circ}$   $+20^{\circ}$   $+60^{\circ}$   $+100^{\circ}$   $\sigma$   $10^{4}$   $20$   $11$   $8$   $54$   $6$   $83$   $5$   $67$   $5$   $28$   $4.54$   $4$   $04$ 

The specific refraction of thallum in its salts is (Gladstone and Dale's formula, Ha line) 0 106, and the atomic refraction is therefore 21 6 14 Thallium and its compounds are diamagnetic (Lamy), the magnetic susceptibility of thallium at the laboratory temperature being -29 x 10 ° cgs electromagnetic units per unit volume 15

In appearance, hardness, clasticity, density, fusibility, specific heat, and electrical conductivity it will be noticed that thallium closely resembles lead. The electromotive behaviour of thalluun is discussed later (p. 173).

<sup>1</sup> M Weiner, Zeitsch. anoig Chem., 1913. 83, 275, cf. Levip, ilad., 1905, 45, 31. Wilhams, ibid., 1906, 50, 127.

2 Richards and Wilson, Conegie Institution Publications, Washington, 1909, No. 118; Zeitsch. physikal. Chem., 1910, 72, 129.

3 Fireau, Compt. rend., 1869, 68, 1125.

4 Richards and others, J. Imer. Chem. Soc., 1909, 31, 154, 1915, 37, 1643.

5 Heycock and Neville, Trans. Chem. Soc., 1894, 65, 31; Peticuko, Zeitsch. anorg. Chem., 1906, 50, 133, see Burgess, J. Washington Acad. Sci., 1911, 1, 15, Chem. News., 1911, 104, 165.

5 Toepler, Wied Annalen, 1891, 53, 343; Pacher, Nuovo Cimento, 1895, [iv.], 2, 143.

7 Robeitson, Trans. Chem. Soc., 1902, 81, 1233, cf. Heycock and Neville, loc. cit.

5 Kiaffs and Knoche, Ber., 1909, 42, 202.

10 Issae and Tammanin, Zeitsch. anorg. Chem., 1907, 55, 58.

10 von Wartenberg, Zeitsch. Elektrochem., 1913, 19, 482, cf. Isiallt and Knoche, loc. cit.

11 Lamy, loc. cit. (p. 165), Regnault, loc. cit. (p. 165), Schmitt, Proc. Roy. Soc., 1903, 72, 177; Richards and Jackson, Zeitsch. physikal. Chem., 1910, 70, 414.

18 Nernst and Schweis. Sitzungsber. K. Akad. Wiss. Berlin, 1914, p. 355.

19 Interpolated from the data of Dewar and Fleming, Phil. Mag., 1893, [v.], 36, 271, and Dickson, tod., 1998, [v.], 45, 525, cf. de la Rive, Compt. rend., 1863, 56, 588, Matthressen and Vogt, Phil. Mag., 1863, [v.], 45, 525, cf. de la Rive, Compt. rend., 1863, 56, 588, Matthressen and Vogt, Phil. Mag., 1863, [v.], 45, 525, cf. de la Rive, Compt. rend., 1863, 56, 588, Matthressen and Vogt, Phil. Mag., 1863, [v.], 45, 525, cf. de la Rive, Compt. rend., 1863, 56, 588, Matthressen and Vogt, Phil. Mag., 1863, [v.], 45, 525, cf. de la Rive, Compt. rend., 1863, 56, 588, Matthressen and Vogt, Phil. Mag., 1863, [v.], 45, 525, cf. de la Rive, Compt. rend., 1863, 56, 588, Matthressen and Vogt, Phil. Mag., 1863, [v.], 45, 525, cf. de la Rive, Compt. rend., 1873, 76, 224.

19 Honda, Ann. Physik, 1910, 32, 1027; Owen, ibid., 1912, [v.], 37, 657.

169 4 THALLIUM.

The flame, are, and spark spectra of thallium are characterised in the visible region by the green ray \$\lambda 5350-7\$, the observation of which led to the discovery of the element by Crookes! The most intense lines in the are and spark spectra are the following (Exner and Haschek) -- 2

aic. 2709:33, 2767 96, 2018:42, 2921:66, 3229 89, 3510:37, 3520:53, 3775'89, 5350'70.

spark: 3519:37, 3529 53, 3775:89, 5350 70.

The most persistent lines in the spark spectrum, and therefore the lines that should be looked for when seeking traces of thallium, are (Evner and Haschek's wave lengths) 2298-25, 2530:94, 2768-00,\* 3091-88, 3519-35,\* 3775-89,\* and 5350 69,\* those asterisked being the most sensitive 3

Commercial thallorm, when spectroscopically examined, is found to contain traces of lead, tin, copper, and aluminium 4

Thallium does not unite with hydrogen 5 With each of the halogens it combines directly. When heated in oxygen, thallium is converted into thallous or thallic oxide, according to circumstances
it at ordinary temperature, producing thallic oxide <sup>6</sup>
Thallium oxideses in an, slowly at the ordinary temperature, rapidly at 100°
The oxidation also occurs under water containing dissolved an, so that thallium is best preserved under an free water or glycerol At a red heat, thallium decomposes water. Thallum combines directly with sulphur, selemum, tellurium, phosphorus, arsenic, and antimony, but not with carbon, silicon, and boron. It is unacted upon by carbon diovide at 300°. Thallium is insoluble in liquid ammonia.

Thalhum dissolves in mineral acids with the formation of thallous salts. The action is very slow with hydrochloric acid, quicker with sulphuric acid, and rapid with intric acid. Except in the cases of intric and concentrated sulphuric acids, hydrogen is evolved. Thallium reduces potassium permanganate at the ordinary temperature

In accordance with its position in the electromotive series (p. 173), thallium readily displaces gold, silver, mercury, copper, and lead from aqueous solutions of their salts.

Thallium is employed in the manufacture of certain kinds of optical glass of high refractive index, since the salts of thallnum are characterised by their unusually high refracting power

Atomic and Molecular Weights. - The atomic weight of thallium is equal to its combining weight in thallons salts, or three times its combining weight in thallic salts. This follows from the isomorphism of (i) numerous - .... ------

<sup>1</sup> The green may is really a triplet (Fabry and Perot, Ann. Chim. Phys., 1899, [vii.],

The green tay is reany a triplet (rang) and room, some field, 115)

2 Kaysei, Handbuch der Spektroskopie (Leipzig, 1900-12), vol. vi. p. 709, where a bibliography will be found; Exner and Haschek, Itie Spektroskoi Elemente ben normalem Prink (Leipzig and Wien, 1911), Huppers, Zeitzeh wiss Photokem, 1913, 13, 16 (atc.).

3 Hartley, Phil. Trans., 1884, 175, 1-325, de Gramont, Compt. send., 1907, 144, 1101, 1910, 151, 308; 1914, 159, 5; Hartley and Moss, Proc. Roy. Soc., 1912, A, 87, 38. For the vacuum tube spectrum of thallum, see Pollok, Sci. Proc. Roy. Dubl. Soc., 1912, 209. For the vacuum time special and the state of 
thallous, potassium, rubidium, and ciesium salts, (ii ) various double thallic halides with the corresponding indiana salts (p 5). It is, moreover, in barmony with Dulong and Petit's Law

The atomic weight of thallium has been determined with great care by Crookes,2 who effected the synthesis of thallons intrate. As the mean of ten extremely concordant experiments, he obtained the following result (0=16 000 , N=11 008) -

TI TINO, 
$$100 \ 130 \ 3910$$
 . TI =  $204 \ 034$ 

The determinations of Lepierre 3 may be looked upon as yielding corroborative evidence in favour of Crookes's value, the results ranging from 201 021 to 204 300 for the atomic weight, and being derived from the analyses of thallous sulphate, thallous intrate, and thallic oxide 4. The value at present accepted for the atomic weight of thallium is Tl = 204.

According to Biltz and Meyer, thallium vapour at about 1700° is diatomic Their results, however, are unreliable, since thallium attacks porcelain, of which their apparatus was made, and von Wartenberg has found, by using an iridium bulb in a Victor Meyer vapour density apparatus, that thallium vapour is essentially monatomic, the molecular weight being about 220 5

In dilute solution in incremy, the molecule of thallium is monatomic, as has been shown by the vapour pressure determinations of Ramsay and of Tammann, and the E M F measurements of Richards and Wilson 6 Moreover, it is monatomic in dilute solution in tin, bismuth, cadmium, zinc, and sodium, as the experimental and cilculated values for the depression of the freezingpoint of the solvent show

|                                             |       | • |    |                                    | ns bong<br>eston of                  | 1 |
|---------------------------------------------|-------|---|----|------------------------------------|--------------------------------------|---|
| . Sol                                       | vent. |   |    | Found                              | Calculated                           | : |
| Tm<br>Bismuth<br>Cadminin<br>Zine<br>Sudiim | :     | : | .' | 2 9<br>2 0'<br>1 5<br>5 0'<br>4 2' | 3 1°<br>2 2°<br>1 6'<br>5 2°<br>1 4° |   |

<sup>1</sup> See Clarks, A Revalentation of the Rome, Weights, 3rd ed. (\* Smithsonian Miscellaneous Callections," vol. 54, No. 3, 1910), p. 273

2 Crookes, Phil. Trans., 1873, 163, 277, Chem. New., 1874, 29, 14, etc.
3 Lepicite, Bull. Size thim., 1893, [iii.], 9, 166, 1894, [iii.], 11, 123, Compt. rend., 1893, 116, 580

4 For other determinations, see Lamy. Inn. Chem. Phys., 1863, [iii.], 67, 410. Helberting, Annalus, 1865, 134, 11, Weither, J. prakt. Chem., 1864, 92, 128, 351; Wells and Penfield, Amir. J. See, 1894, [iii.], 47, 166

5 H. Bitt and V. Meyer, Br., 1889, 22, 725, 11. Bilty, Zeitsch. physikal. Chem., 1896, 19, 385, von Wartenberg, Zeitsch. anorg. Chem., 1997, 56, 320

4 Ramsey, Trans. Chem. Soc., 1889, 55, 521, Tammann, Zeitsch. physikal. Chem., 1889, 3441, Rehards and Wilson, dud., 1910, 72, 129, Carnegoe Institution Publications, 1909, No. 118. It is more correct to say that the molecules Tillige exist in a dilute solution of thallium in mercury (J. Hiddebrand, J. Amir. Chem. Soc., 1913, 35, 501, J. Hiddebrand and E. D. Eastman, dod., 1915, 72, 2452, M. Phall Smith, dud., 1914, 36, 847, Beckmann and Luesche, Zeitsch. anorg. Chem., 1914, 89, 171)

7 Heycock and Neville, Trans. Chem. Soc., 1889, 55, 666; 1890, 57, 376, 1892, 61, 888; 1897, 71, 383. The experimental numbers represent the depression produced by

## COMPOUNDS OF THALLIUM,

General, - Thallom forms two series of salts thallows salts (TIA1), derived from the powerfully basic oxide thallows oxide (TLO), and thallie salts (TIV), derived from the weakly basic oxide thallie oxide (TLO) - All the salts of thalloun are poisonous, producing symptoms like those of lead poisoning 1

Thallous compounds and the compounds of the alkali metals have many points of similarity. The salts derived from colourless acids are generally colourless, and those which are soluble in water crystallise readily, usually as anhydrous salts. They are volatile at a red heat. Many soluble thallous salts are isomorphous with the corresponding salts of ammornin, potassium, or rubidium, a fact that was early recognised  $^2$  . Thallous hydroxide is a strong base. It is readily soluble in water, forming a solution which readily absorbs carbon dioxide, is soapy to the touch and alkaline in reaction

Both in appearance and in solubility, a number of sparingly soluble thallous salts (chloride, brounde, iodide, sulphide, chromate, etc.) closely resemble the corresponding salts of lead

In aqueous solution the thallous salts are ionised to approximately the same extent as the corresponding salts of the alkali metals. The following values for the percentage dissociation at 18' are given by A. A. Noves and

| Nomolny                                                  | 001                  | 002   001                                      | 0 01                         | 0 02   0 05                    | 0 1          | 0 2  |
|----------------------------------------------------------|----------------------|------------------------------------------------|------------------------------|--------------------------------|--------------|------|
| Tic1<br>111b<br>111NO<br>TI <sub>2</sub> SO <sub>4</sub> | 97 6<br>97 7<br>91 9 | 96.5 94.2<br>96.1<br>96.7 (91.8<br>92.4 (88.2) | 91 5<br>92 6<br>92 6<br>93 7 | 90 8 86 5<br>91 3<br>78 0 69 1 | 78 8<br>62 5 | 56 1 |

I grain around thellium in 100 grain froms of olvert, the calculated numbers are those drived from Van't Holl's formula (Vol. 1 p. 146) for 1 gram molecule of solute in 100 grams of solvent and edemated to 100 gram atoms of solvent. For sedimin, E. Griffiths' value for the latent heat of fission of sodium, 27.5 c. its per grun, was used (Griffiths, Proc. Roy,

of solvent and almated to 100 gram atoms of solvent. For schom, E. Griffiths, value for the latent heat of themon of sodium, 27.5 cals per gram, was used (Griffiths, Proc. Roy. No. , 1911, A, 89, 561).

Lamy, Compt. rend., 1863, 57, 412. Paulet, chid., 1863, 57, 494, Blake, chid., 1899, 111, 57, (unc), Chong rend., 1863, 57, 412. Paulet, chid., 1863, 57, 494, Blake, chid., 1899, 111, 57, (unc), Chong Zente, 1895, p. 33, 1898, p. 120.

The literature on the isomorphism of thallom salts is vely extensive. See de la Provostay, Compt. rend., 1862, 55, 610. von Lang, Phil. May, 1863, [v.], 25, 218, Muller, Proc. Roy. Soc., 1865, 144, 555, Morther, J. prick Chong, 1866, [v.], 25, 218, Muller, Proc. Roy. Soc., 1866, 19, 594, Ramma Isberg, Ber., 1870, 3, 276, Progr. London, 1872, 146, 552, Schreder, Ber., 1874, 7, 676. Fock, Actich Kryd, Mar., 1884, 49, 1897, 28, 337. Wyroubedl, Bull. See Gram, Mar., 1884, 25, 32, 1884, 7, 139. Doket Compt., cond., 1884, 99, 807, Soc., told., 1884, 99, 390, Lepueria and Lachand, that., 1891, 113, 196, Retgers, Fetch. physikal Chem., 1889, 4, 623, 1891, 8, 13, Royerboun, that., 1891, 1, 22, Gessner, Edd., 1899, 30, 430, 1995, 54, 771, Proc. K. Mad., Hernsch, 1899, 1, 224, Gessner, Edd., 1899, 30, 430, 1995, 54, 771, Proc. K. Mad., Hernsch, 1899, 1, 224, Gessner, Edd., 1894, Mir., 1903, 38, 110, Storiebleke, Rec. trac. chem., 1902, 21, 87, 1905, 24, 53, 1907, 26, 245, Wells and Pentheld, Lappond, Chem. Nov., 1893, 74, 247, Tattion, Proc. Roy. Soc., 1907, A, 79, 331, 1910, A, 83, 211, Wallace, Zeiteck Krist, Mar., 1911, 49, 417, Rosenbasch, ibid., 1909, 33, 99; Walleraut, Bull. Soc. Grang. Mar., 1905, 28, 311.

At 18° the ionic mobility of the thallous ion is 65.55,1 and, adopting Kohlrausch's value for its temperature-coefficient, the ionic mobility at 25° is 75 48 The corresponding value for potassium is 64 7 at 18°. The transport number for the thallous ion in dilute solution is 0.479 at 25°2 The equivalent conductivities at 25° of a number of thallous salts are tabulated below -3

| v -                                                                        | 4 8         | 16    | 32                                                             | 61                                                                      | 128                                                                                    | 256                                                                                    | 512                                                                                             | 1021                                                                          |
|----------------------------------------------------------------------------|-------------|-------|----------------------------------------------------------------|-------------------------------------------------------------------------|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------|
|                                                                            | - '         | ''    |                                                                |                                                                         |                                                                                        |                                                                                        |                                                                                                 |                                                                               |
| TIOH TIF TICI TICIO, TICIO, TIGIO, TISO, TISO, TI,SO, TI,SO, TI,SO, TI,SO, | 182 0 200 0 | 217 0 | 230·0<br>115·9<br>123·6<br>129·3<br><br>113·1<br>83·0<br>111·2 | 238·0<br>120 6<br><br>127·8<br>134 0<br>122 9<br>141 9<br>94·9<br>120 7 | 241.0<br>123.7<br>139.6<br>129.8<br>137.5<br>122.9<br>131.2<br>151.7<br>106.1<br>129.0 | 248 0<br>126 2<br>143 1<br>132 1<br>139 6<br>125 5<br>138 3<br>160 2<br>115 2<br>134 7 | 248 0<br>128·1<br>145·1<br>134·2<br>141 9<br>126 3<br>111 5<br>143 1<br>166·7<br>123·7<br>138 6 | 130·1<br>135·4<br>143 7<br>128 1<br>112 0<br>146·4<br>170 6<br>130 8<br>142·2 |
| TINO,<br>TIH, 1'O,<br>TI, 2HA, O,<br>TI, 2CO,<br>TI, 2CQ,                  |             | 70 4  | 74 3<br>93·5<br>105 0                                          | 133 8<br>96 9<br>78 3<br>107 3<br>118 0                                 | 137·6<br>101 1<br>81·2<br>119 2<br>128 5                                               | 101·1<br>104 0<br>82 8<br>129 9<br>139 0                                               | 142.0<br>106.5<br>84.2<br>137.1<br>117.7                                                        | 142.6<br>108.7<br>85.4<br>143.4<br>153.5                                      |

Numerous investigations have been carried out dealing with the influence of other salts on the solubilities of thallous salts. The results may be briefly summarised as follows. For uni-univalent thallous salts the changes of solubility caused by other salts are in qualitative agreement with the deductions from the ionic theory and the law of chemical equilibrum, and quantitative agreement, though never exact, is approached most closely with the least soluble thallous salts For umbivalent thallous salts, the solubilities are affected by salts having a common univalent ion (i.e. the thallous ion) in a similar manner; but the influences exerted by other salts, having bivalent ions in common with the thallous salts, are not even in approximate agreement with theoretical deductions unless it be assumed that the umbivalent salts undergo dissociation in two stages, eq. Tl<sub>2</sub>SO<sub>4</sub> = Tl' + TlSO<sub>4</sub>', TlSO<sub>4</sub>' = Tl + SO<sub>4</sub>", and that even in dilute solution the concentrations of the intermediate ions are considerable 4

Thallous hydroxide is considerably less stable than an alkali hydroxide

Bates, J. Amer. Chem. Soc., 1913, 35, 519; cf Kohliausch, Zeitsch physikal. Chem., 1905, 51, 741.
 Falk, J. Amer. Chem. Soc., 1910, 32, 1555.
 Franke, Zeitsch physikal Chem., 1895, 16, 463, see also A. A. Noyes, Farrell, and Stewart, vide info. Bay and Winninghoff, inde info., Hunt, J. Imer. Chem. Soc., 1911, 33, 1650; Kohliausch and Steinwicht, Sitzungsber. K. Akad. Wiss. Berlin, 1902, p. 581, Kohliausch and Maltby, ibid., 1899, p. 665. In the table, v=dilution, in littes per gramaguraluri.

 <sup>4</sup> A. A. Noyes, Fattell, and Stewart, J. Amer. Chem. Soc., 1911, 33, 1650, Biay and Winninghoff, ibid., 1911, 33, 1663, Biay, ibid., 1911, 33, 1673; Harkins, ibid., 1911, 33, 1886, see also Aberg and Spencer, Zeitsch. among. Chem., 1905, 46, 106; A. A. Noyes, Zeitsch. physikal. Chem., 1890, 6, 241; 1892, 9, 603; A. A. Noyes and Abbot, ibid., 1895, 46, 129.

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as is indicated by its relatively small heat of formation. It is easily dehydrated (even at 100°) to thallous oxide, which has a very small heat of hydration. Further, thallium is considerably less electropositive than the alkali metals. According to the best experimental results available (1916), the following sequence of metals in the electromotive series is correct and indicates the position of thallium

. . Zn, Cd, Fe, Tl, Co, Sn, N1, Pb . . .

Thallium has therefore a decided tendency to assume the iome state. The electrode potential of thallium at 25° is -0.6170 volt (normal calomel electrode being zero) For very dilute solutions, the potential of the thallium electrode varies with the concentration of thallous ion in the liquid bathing it in strict accordance with the Nernst formula, a slight deviation is noticeable when the concentration of thallium ion reaches 0.1 normal.1 The accuracy with which the Neinst formula is capable of expressing the results is strong evidence against the hypothesis of the existence of subvalent thallmin ion Tl  $_{\mathcal{B}}$  which has been supposed by Denham  $^2$  to exist

Thallic compounds resemble the compounds of aluminium to a certain extent. The thallic salts are in general readily soluble in water, and crystallise with considerable amounts of water of crystallisation. They are derived from a very weak base, thallic hydroxide, which is practically insoluble in water, and which in case of dehydration resembles aurie hydroxide. The thallie salts are therefore considerably hydrolysed by water, and solutions of these salts are only stable in the presence of an excess of acid. A solution of thallic sulphate, for example, containing a slight excess of sulphuric acid, gives a brown precipitate of thallic hydroxide when diluted or warmed. Thallic salts are decidedly unstable, anhydrous thallic chloride, for instance, losing chlorine at temperatures below 100°, and they exhibit a great tendency to form complex salts

Thallous salts may be exidised to thallie salts and rice versa, the former conversion being not so readily accomplished as the latter. The reduction of thallie to thallous salts may be readily and quantitatively effected by the ordinary reducing agents, and it therefore happens that the addition of ammonium sulphide to a solution of a thallous salt leads to the precipitation of thallous sulphide and sulphur Further, potassium undide gives a precipitate of thallous include and rodine when added to a thallic salt salts are quantitatively reduced to thallous salts by thallium itself. Thallous salts may be oxidised by potassium permanganate, the reaction when effected in the presence of hydrochloric acid and under certain conditions being sufficiently exact to be used for the estimation of thallium. Practically complete oxidation of thallous salts can also be brought about by chlorme or

Thallous and thallic salts exhibit to a high degree the curious property of combining with each other to form what may be termed intermediate salts such, for example, as the chlorides of the composition Tl<sub>2</sub>Cl<sub>3</sub>(TlCl<sub>3</sub>, 3TlCl) and

G. N. Lewis and Van Ende, J. Amer. Chem. Soc., 1910, 32, 732.
 cf. Neumann, : Zeitsch physikal Chem., 1891, 14, 193, Spencer, ibid., 1911, 76, 360.
 Abegg and Spencer, Zeitsch anorg. Chem., 1905, 46, 406.
 Shukoff, Ber., 1905, 38, 2691.
 Birslee, Trans.
 Faraday Soc., 1909, 4, 159.
 Denham, Trans Chem. Soc., 1908, 93, 833
 Abegg and Spencer, Zeitsch. anorg. Chem., 1905, 44, 379.

TICl<sub>2</sub>(TICl<sub>3</sub> TICl). Since these salts usually resemble the thallous salts in being sparingly soluble in water, it often happens that such intermediate salts are pro luced during the oxidation of thallous, or the reduction of thallic salts.

The modus operands of these processes, however, is not definitely known. Thermochemistry of Thallium Compounds. —A number of heats of formation, etc , are given in the following table --

| Comp                                         | hund                                    | - | Heat of<br>Formation | Heat of<br>Solution                                                                                                |                                            |
|----------------------------------------------|-----------------------------------------|---|----------------------|--------------------------------------------------------------------------------------------------------------------|--------------------------------------------|
| TIF FIG                                      | : : : : : : : : : : : : : : : : : : : : |   |                      | Cals<br>+51 4 4<br>48 6<br>41 4<br>30 2<br>12 2<br>56 9<br>21 7<br>17 7<br>12 2<br>58 1<br>221 0<br>80 8<br>56 5 8 | Cals -10 1 3 1 - 3 15 10 0 8 3 1 8 1 - 2 1 |
| П,<br>ПСПВг <sub>2</sub> , (П <sub>в</sub> О |                                         | • |                      | 10.8 3                                                                                                             | 2 2<br>- 2 9                               |

```
0, H + [0, T]
                      -2[TIOH] + 3.25 Cals
                     = TlCl aq + 13.71 ,,
   ,,
TIOHaq. +\frac{1}{2}H<sub>2</sub>SO<sub>1</sub> aq =\frac{1}{2}TISO<sub>1</sub> aq +15 6
```

Alloys.4 -Thallium readily forms alloys with many other metals - The

<sup>1</sup> See V Thomas, Ann Chim Phys., 1907, [vin.], 11, 204.

2 J. Thomsen, Popy Annalin, 1871, 143, 354, 197, J. publi, them., 1875, [n.], 12 98, 1879, 19, 13, 1880, 21, 38, 419. Tommasi, Compt. rend., 1884, 98, 812. Petersin, Zelish physikal Chem., 1889, 4, 381. Fabre, Ann. Chim Phys., 1887, [vi.], 10, 172, 1885, [vi.], 14, 115. Thomas, that, 1907, [vin.], 11, 204.

3 Including the heat of solution of the compound.

4 With K, Carstangen, J. prait Chem., 1897, 102, 65. Heycock and Neville, Trans Chem. Soc., 1889, 55. 666. Biedig and Haber, Rev., 1898, 31, 2741. Kunnakow and Puschin, Zeitsch. anorg Chem., 1902, 30, 86. with Na. vide pieceding references and Tammann, Zeitsch. physikal Chem., 1899, 3, 441. with Cu. Carstanjen, low cit., Documekel Zeitsch. anorg Chem., 1906, 48, 185. with Ag, Heycock and Neville, Trans Chem. Soc., 1894, 65, 31. Osmond and Robeits Austen, Phil. Trans, 1897, A, 189, 25. Petrenko, Zeitsch. anorg Chem., 1906, 50, 183. with Phil. Trans, 1896, A, 187, 417. Robeits-Austen, ibid., 1888, 179, 339. Levin, Zeitsch. anorg. Chem., 1905, 45, 31. with Ca, Donski, ibid., 1908, 57, 185. Ban, ibid., 1911, 70, anorg. Chem., 1905, 46, 76, with Zu. Carstanjen, loc. cit. Heycock and Puschin. Trans. Chem., 1907, 52, 30. with Cd. Carstanjen, loc. cit., Kunakoff and Puschin, Zeitsch. anorg. Chem., 1902, 30, 86; with Hg, Carstanjen, loc. cit., Kunakoff and Puschin, Zeitsch. anorg. Chem., 1902, 30, 86; with Hg, Carstanjen, loc. cit., Kunakoff and Puschin, Zeitsch. anorg. Chem., 1902, 30, 86; with Hg, Carstanjen, loc. cit., Regnauld, Compt. rend., 1867, 64, 611, Humphreys, Trans. Chem. Soc., 1896, 69, 1679, Puschin, J. Russ. Phys. Chem.

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following inter-metallic compounds have been described, containing thallium and a metal of Group 1 or II -

| NaTl,  | աթ | 306  | Tl <sub>3</sub> Mg <sub>8</sub> , mp 413° | CaTl, no m p     |
|--------|----|------|-------------------------------------------|------------------|
| KTI,   | ,, | 335° | TIMg., no m p                             | CaTl, in p. 969° |
| Hg,Tl, | ,, | 15°  | $Tl_{x}Mg_{y}$ ,                          | Ca,TL, no m p    |

Thallum is miscible with gold, silver, and culminm, and with silver forms anixed crystals containing 0 to 10 per cent of thallowing It is only partially

iniscible with copper and zinc, and is practically non-miscible with aluminium. Thallium readily dissolves in mercury  $\Lambda$  liquid analgam containing 8.5 per cent of thallmin may be used in thermometers down to - 60°1.

Alloys of thallum and the metals of Groups IV to VIII in the Periodic Table are mentioned in the corresponding volumes of this series

#### THAILIUM AND THE FILORING GROUP

THALLOUS HALIDES Thallons fluoride, chloride, bromide, and iodido are all known. The last three salts resemble the corresponding compounds of lead very closely in appearance and in physical properties, it has been both affirmed and denied that in the liquid state they have the properties of liquid crystals.2 The molten chloride and brounde have not been obtained optically clear

Thallons chloride, bromide, and iodide are miscible in the liquid state, and the three systems TICL TIBE, TIBE TH, and TICL—TIL have been studied by the thermal method.—The results show that the first two systems yield contianous series of mixed crystals, the "highdus" curves exhibiting minima, and that the last system forms an incomplete series, the "liquidus" exhibiting a transition point 1

Thallous fluoride, TIF, is prepared by dissolving thallous hydroxide, carbonate, or sulphide in aqueous hydrofluoric acid, evaporating to dryness, heating to remove hydrogen fluoride, and crystallising the residue from water. It may also be prepared by heating thallous oxide or carbonate to dull redness in a stream of dry hydrogen fluoride. Thallium dissolves slowly in aqueous hydroflooric acid

Thallous fluoride is readily soluble in water, from which it crystallises in colourless, lustrous, regular octahedra. At a red heat it may be distilled in a current of hydrogen fluoride. It is insoluble in alcohol and in liquid

ammonia. The aqueous solution has an acid reaction. The monochure crystals and to separate in monochure crystals (1,0) is said to separate in monochure crystals. from a solution of thallous carbonate in aqueous hydrofluoric acid s and

Suchun, Zeisch Licktockem, 1906, 12, 72c, T.W. Richards and Pusclan, loc. et.; Suchun, Zeisch Licktockem, 1906, 12, 72c, T.W. Richards and Wilson, Zeisch physikal. (Term., 1910, 72, 12c), J. Hiddeland, J. Imr. Chem., Soc., 1913, 35, 501, with Al, Carstanger, loc. etc., Doernekel loc. ed.

1. M. Intosh and Johnson, J. Imr., Chem. Soc., 1912, 34, 910.

2. Stoltzenberg and Huth, Zeisch physikal Chem., 1910, 71, 641, Tabanelt and E. Lorenz, Festscheif W. Kenns 1912, p. 446

2. R. Lorenz and Estel, Zeisch among Chem., 1915, 91, 61.

4. Monkemeyer Johnh. Min. Bert., 1996, 22, 1

5. Buchner, J. prof.t. Chem., 1805, 95, 404, Sitzungsber K. Akad. Wiss. Wien, 1865, [11, 52, 041]. Epinam and Bartecko Zeisch among Chem., 1909, 61, 238.

5. Kublimann, Compt. rend. 1864, 58, 1036.

7. Gore, Pro. Roy. Soc., 1873, 21, 140.

8. Kublimann, loc. ed., Willin, Ann. Chim. Phys., 1865, [1v.], 5, 5.

thallium hydrogen fluoride, TIF.HF, crystallisos out in regular octahedra when a solution of thallous fluoride in concentrated aqueous hydrofluoric acid is evaporated over sulphuric acid. The acid fluoride is stable at 100°, but decomposes at higher temperatures (Buchner).

Aluminium thallous fluoride, 2AIF, 3TIF, and numerous other double luorides have been prepared.1

Thallous chloride, TICI, may be obtained by the action of chlorine on thallnum. It is prepared by adding hydrochloric acid or a solution of a soluble chloude to a solution of a thallous salt, and forms a white, crystalline precipitate

Thallous chlorade is a white, crystalline solid of density 7.02.2 It melts to a yellow liquid at 426 °C., 3 and boils at 708 ° to 731 °C.4 Its compressibility coefficient is 48 × 10 <sup>-6</sup> per atmosphere <sup>5</sup> The chlorade crystallises in isometric cubes and is isomorphous with potassium chloride 6 From 830° to 1030° its vapour density is in agreement with the molecular formula TiCl,7 and the molecular formula is also TCI in solution in fused mercuric chloride.

Thallous chlorde is slightly soluble in cold, and more readily soluble in hot water. The solubility, in grams per litre, is as follows .- 9

Temperature, °C. 0" 10° 20° 25° 30° 40° 50° 60° 80° 100° Grams of TiCl 17 24 34 40 46 60 80 102 160 241 Grams of TICI

The chlorde is less soluble in dilute solutions of hydrochloric acid and alkalı chlorides than in pure water, it is insoluble in alcohol The influence of various salts on the solubility of thallous chloride in water at 25 is shown by the following table of results, due to Bray and Winninghoff, solubilities being expressed in milli equivalents per litre - 10

| Conc of<br>Added<br>Salt in              |                              |                              |                                       | -<br>Added Sal                | <br>t.                             |                                           |                                           |
|------------------------------------------|------------------------------|------------------------------|---------------------------------------|-------------------------------|------------------------------------|-------------------------------------------|-------------------------------------------|
| milli-eqs<br>per litie                   | HCI.                         | KC1.                         | BaCl <sub>2</sub>                     | TINO,                         | Tl <sub>2</sub> SO <sub>4</sub>    | KNO <sub>3</sub>                          | K <sub>2</sub> SO <sub>4</sub>            |
| 0<br>20<br>25<br>50<br>100<br>200<br>800 | 8 66<br>5 83<br>3 83<br>2 53 | 8.69<br>5.90<br>3.96<br>2.68 | 16 07<br>8 98<br>6 18<br>4 16<br>2 82 | 16 07<br>8 80<br>6 24<br>4 22 | 16 07<br>10:31<br><br>6 77<br>1 68 | 16.07<br>17.16<br>18.26<br>19.61<br>23.13 | 16 07<br>17:79<br>19 42<br>21 37<br>26 00 |

<sup>1</sup> Ephnam and Barteczko, loc cit, and subsequent volumes in this series.
2 Lamy, Ann. Chim. Phys., 1803, [iii], 67, 385. F. W. Clarke, Ann. Chim. 1883, 5, 240.
3 Monkomeyer, Jahrb. Min Bed., 1906, 22, 1, Camelloy (Trans Chem Soc., 1878, 33, 273) gives 427°, Stollzenberg and Huth (Zeitsch. physikal. Chem., 1910, 71, 641) give 407°, and Sandomini (Alt. R. Accad. Lincer, 1915, [v.], 24, 1, 616) gives 129°.
4 Carnelley and Williams, Trans. Chem. Soc., 1878, 33, 281.
5 Richards and Jones, J. Amer. Chem. Soc., 1878, 33, 281.
6 Stortenbeker, Rec trav. chim., 1905, 24, 53.
7 Roseco, Proc Roy. Soc., 1878, 27, 426. Ber., 1878, 11, 1196.
8 Beckmann, Zeitsch. anory Chem., 1907, 55, 175.
9 Earl of Berkeley, Phil. Trans., 1904, A, 203, 208., A. A. Noyes, Zeitsch. physikal., 26m., 1892, 9, 603; Bottger, ibid., 1903, 46, 602, Kohliausch, ibid., 1903, 44, 197, 1904, 50, 855, Hill and Siminons, J. Amer. Chem. Soc., 1909, 31, 821.
18 Bray and Winninghoft, J. Amer. Chem. Soc., 1911, 33, 1663

. The specific conductivity of the saturated aqueous solution is  $1514\times 10^{-8}$ reciprocal ohms per cm cube at 18°, and 2176 10-6 at 25° C 1

Fused thallous chloride is miscible with many other molten chlorides, and from a study of the freezing-point curves it has been shown that various double chlorides exist, a number of which are stable at the melting-point. The compounds at present known containing thallium and the metals of Groups I and II, are as follows -2

```
2AgCl 3TlCl
                               TICI 2ZuCl<sub>2</sub> (m p 226')
Ticl ('aCl, (m p 683°)
                               TICI CdCl, (mp. 436)
TICLS(CL)
                               TICI HgCl, (m.p. 221')
TICL 2MgCL
                               4TICI HgCl,
2TICl ZnCl<sub>2</sub> (m p 352')
```

Thallous chloride forms mixed crystals with the chlorides of potassium, rubidium, zinc, and mercury — The double salt TICLHgCl $_2$  crystallises in needles, and was originally prepared by Carstanjen,3 who also prepared thallous an whiler ide, TI AuCI.

Thallous bromide, TBr, is prepared like the corresponding chloride, It is a pale yellow solid of density 7.54.1 and its compressibility coefficient is 5.2 - 10 ° per atmosphere 5 - It crystallises in the isometric system in cubes, melts to a brownsh-vellow liquid at 450, and boils at 800 to 811 7. The brounde is stable at high temperatures  $^{8}$ 

The solubility of thallous brounde in water, in grains per 100 grains of water, is as follows -9

The specific conductivity of the saturated aqueous solution is 192 10-6 reciprocal ohms per cm cube at 18 10

Thallous iodide, TH, is prepared like the corresponding chloride and brounde, and forms a citron yellow precipitate of density 7 07.11 crystalline form is orthorhombic. At 168, the yellow thallous iodide is converted into a red, cubic variety, 12 the compressibility of which is  $6.8 \times 10^{-6}$ 

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<sup>1</sup> Kohlrausch, Zeitsch physikal Chem. 1903, 44, 197, 1904, 50, 355, Bray and
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<sup>\*\*</sup>Nonmansin, Science (1997) \*\*Nonmansin, Science (1997) \*\*\*Inches (1997) \*

<sup>6, 82.</sup>Lamy, Ann. Chim Phys., 1863, [m.], 67, 385, F. W. Clarke, Amer. Chem. J., 1883,

per atmosphere,1 and the density 7.10. The two polymorphs are interconvertible. The red form melts at 431° to a dark red liquid 2

The conversion of one form into the other is often extremely slow. A hot, filtered, aqueous solution of thallous iodide deposits red crystals when rapidly cooled, but a hot, saturated solution, when slowly cooled in the presence of the yellow form, deposits yellow crystals. A hot solution in aqueous potassium acetate or hydroxide deposits the metastable red modification when cooled, and the red form may also be precipitated by adding potassium iodide to a hot, concentrated solution of a thallous salt 3. Freshly precipitated yellow thallous inclide becomes green when exposed to similaht. The green holdo can be recrystallised from water without losing its colour, but on standing it slowly reverts to the yellow modification, the change being hastened by heating with aqueous iodine or potassium iodide 1

The solubility of the yellow modification in water, in grams per 100 grams of water, is as follows -

```
Temperature, C.
                   0,
                            20"
                                    10'
  601
  80
   100
                 0.002
                         0.006
Grams of TH
                                  0.015
   0.035
   0.070
  0.120
```

The specific conductivity of the saturated aqueous solution at 18/18/22/3 - 10<sup>-6</sup> recipiocal ohms per cm cube 6

Thallous iochde forms mixed crystals with thallous nitrate, the "hquidus" curve for the system Til - TINO, consists of two parts meeting at a transition point.7

THALLIC HALIDES - Thallic fluoride, TIF, is not known, but unstable, white crystals of thallic potassium thuoride, 2TIF KF, have been

Thallic fluochloride, TIFCl, 3H2O, is obtained in white needles by the action of chlorine on thallous fluoride dissolved in aqueous hydrofluoric acid. It becomes anhydrous when kept over phosphoric anhydride, and the anhydrous salt forms a double salt with potassium chloride, of the formula 2TIFCI, KCI, and a compound with ammonia, of the formula TIFCI, 4NII, Analogous compounds of the composition TIFBr, and TIFBr, 1NH, are also

Thallium oxyfluoride, TIOF, is obtained as an obvergreen powder, insoluble in water, by the action of cold, aqueous hydrofluoric acid on thallic oxide 9

Richards and Jones, J. Amer. Chem. Soc., 1909, 31, 158
 Monkemeyer, Jahrh. Min. Berl., 1906, 22, 1, Carnelley (Trans. Chem. Soc., 1878, 33, 273) gives 439°, Stoltzenberg and Huth (Zeitsch. physikal. Chem., 1910, 71, 641) give 422°, and Van. Eyk. (Proc. K. Akad. Wetersch. Amsterdam, 1901, 3, 98) gives 422°.
 Gerner, loc. etc., and Compt. rend., 1904, 139, 278, 1909, 148, 1015. Ann. Chem. Phys., 1909, [vin.], 17, 290. Willin, ibid., 1865, [iv.], 5, 5, Hebberling, Annalen., 1865, 134, 11. Weither, J. prake Chem., 1864, 92, 128.
 Knosel, Ber., 1871, 7, 576, 803
 Bottger, Zeitsch. physikal. Chem., 1903, 46, 602, Kohlausch, ibid., 1903, 44, 197;
 1901, 50, 355; Long, Zeitsch. anal. Chem., 1903, 36.
 Kohlrausch, ibe. etc.
 Van. Eyk., Proc. K. Akad., Wetersch. Amsterdam, 1901, 3, 98.
 Gewecke, Annalen, 1909, 366, 217.
 Gewecke, loc. etc.; ef. Willin, Ann. Chem. Phys., 1865, [iv.], 5, 5.

Thallic chloride, TICl, -- Thallic chloride cannot be obtained in the pure state by the chlormation of thallons chloride except when the reaction is carried out in a scaled tube and the pressure of the chlorine is several atmospheres. Thallie chloride thus prepared is described by Thomas as consisting of white crystals, which melt at 60 to 70' (iii a

Anhydrous thallie chloride may be obtained by dehydrating its tetrahydrate at the ordinary temperature over sulphume acid, potash, or phosphoric anhydride, or, according to Meyer, by the decomposition in raino of the compound TiCl (C.H.).O The chloride thus obtained forms small, six-sided plates, and melts at 25 — It commences to lose chlorine at 40", and the decomposition is rapid at 100'. It is extremely hygroscopic, rapidly absorbing moisture with the formation of an aqueous solution of thallie chloride, and it is soluble in alcohol, ether, and other organic solvents. With a small quantity of water, thallie chloride forms a clear solution, which is considerably hydrolysed by dilution

When thallous chloride is suspended in water and chlorine is passed through the liquid for a sufficient length of time, a clear solution of buillic chloride is obtained. If the solution be evaporated below 60 to a syrup, and then cooled in a freezing mixture, colourless, orthorhombic (t) crystals of the tetrahydrate, TIC1, 4H2O, separate This hydrate melts at 43 to 45°, and m dry an slowly effloresces (more rapidly at 55) to the monohydrate, TICL H<sub>2</sub>O<sub>5</sub> while according to M<sup>\*</sup>Clenah in, a dihydrate, TICL 2H<sub>2</sub>O also The monohydrate crystallises from a saturated aqueous solution of the trichloride at ordinary temperatures.

From a solution of thallie chloride acidified with intric acid, silver intrate quantitatively precipitates the chlorine as silver chloride, but from a neutral s solution, in addition to the precipitate of silver chloride, a brown precipitate of thallic hydroxide is obtained.

Thallie chloride combines we'r ammonia, producing a white, crystalline compound, TiCl, 3NH, decomposed by water. This compound was prepared William a number of ways According to Thomas, a compound, TICL 5XII, as produced by the action of an excess of ammonia quantifydrons thallic chloride. No compound richer in ammonia is known, and the compound TICl, 5NH, is insoluble in liquid animonia. Thallie chloride combines with other, pyridine, andine, etc

Thallic chloride tetrahydrate readily absorbs one molecular proportion of hydrogen chloride, a liquid being produced from which, on standing over phosphoric anhydrade, hydrogen thallie chloride, TiCl, HCL3H2O or HTICl, 3H2O, separates in deliquescent crystals

Thallic bromide, 2 TiBr , is not known in the anhydrous state. It can be obtained in solution by the action of excess of bromine on thallous bromide suspended in water. When the solution is concentrated at 30' to  $40^{\circ}$ to a syrup, and then strongly cooled, very pale vellow crystals of the tetra-hydrate, TBr, 4H,O, are produced. Meyer has obtained the monohydrate,

240

<sup>Willim, Ann. Chem. Phore, 1865, [iv], 5, 5; Weither, J. makt. Chem., 1864, 91, 385.
R. J. Meyer, Zeitech. among. Chem., 1900, 24, 321; 1902, 32, 72. Rev., 1902, 35, 4119.
V. Thomas, Compt. rend., 1902, 134, 545, 135, 1051. Inn. Chem. Phys., 1907, 2011.
J. H., 204. Cushman, Amer. Chem. J., 1900, 24, 222, 1901, 26, 505.
M. Clenahan, Amer. Chem. J., 1904, [iv.], 18, 101.
Renz, Ber., 1902, 25, 1110.</sup> 1902, 35, 1110.

2 Willim, loc. cit., R. J. Meyer, loc cit., Thomas, loc. cit t.

TIBr<sub>s</sub>. H<sub>2</sub>O, by concentrating the solution in vacuo at the ordinary temperature. The hydrates cannot be dehydrated without loss of bromine They absorb one molecular proportion of hydrogen brounde or hydrogen chloride

has described the addition compound, TIBr, 3NH<sub>2</sub>.

Thallic chlorobromides, TIClBr, 4H<sub>2</sub>O and TICl<sub>2</sub>Br, 4H<sub>2</sub>O, have been prepared by Thomas by the action of bromine on thallous chloride suspended in water, and of chlorine on thallous bromide suspended in water, the solutions being concentrated to a syrupy consistency at 30° to 40°, and then strongly cooled The hydrated salts cannot be dehydrated without loss of halogens occurring. The hydrates combine with one molecular proportion of hydrogen chloude or brounde, but the compounds formed have not been obtained in the crystallme state 1

Thallic iodide, TII, may be prepared by digesting thallous iodide with an alcoholic solution of iodine and allowing the solution to evaporate over sulphune acid. It forms large, dark, orthorhombic crystals (a b.c= 0.6828 1 11217), isomorphous with rubidium and caesium til-iodides.2 It might therefore be concluded that thallie iodide is a per iodide of thallous iodide and essentially a thallous compound. According to Abegg and Maitland, it is a tautomeric substance, behaving as a thallous or as a thallic compound according to circumstances. In its sparing solubility in water and organic solvents it resembles the thallons halides, while it resembles the other thallie halides in forming complex salts with other iodides. In aqueous solution these double salts yield the stable complex ion  $\mathrm{TH}_i$ 

Double Salts containing Thallic Chloride, Bromule, or Iodide - A very large number of these double salts is known,

The following compounds with the alkali metal and ammonium halides have been described -

```
Type TIX, MX xH,O.
   Type 2TlX , 3MX J H2O.
 TlBr_3 NH_1Br
  2TICL 3CsCl
TlBr<sub>1</sub>,Nll<sub>1</sub>Br 2H<sub>2</sub>O*
TlBr<sub>1</sub>,NH<sub>1</sub>Br 4H<sub>2</sub>O
  2TlBr<sub>3</sub> 3KBr.3H<sub>2</sub>O
2TlBr<sub>3</sub> 3CsB<sub>1</sub>
2TlI<sub>3</sub>.3K1.3H<sub>2</sub>O
TIBr<sub>3</sub>.NH<sub>4</sub>B<sub>1</sub>.5H<sub>2</sub>O
TIBr, KBr.2H,O*
TIBr, RbBr H,O*
TlBr, CsBr*Tll, NH4I
TH<sub>3</sub>.KI
TH<sub>3</sub>.Csl*
TH<sub>3</sub>.RbI 2H<sub>2</sub>O*
```

<sup>&</sup>lt;sup>4</sup> V. Thomas, Compt read., 1902, 135, 545, 1051, 1208, Ann. Chim Phys., 1907, [viii.], 11, 201.

Wells and Penfield, Amer J. Sci., 1894, [iii.], 47, 463, Zedsch anora Chem., 1894, 6, 312, cf. Nickès, Compt. rend., 1861, 58, 537, J. Pharm Chim., 1864, [iv.],

<sup>1894, 6, 312,</sup> cf. Nickès, Compt. rend, 1861, 58, 537, J. Pharm Chim., 1861, [iv.], 1, 25.

3 Abegg and Maitland, Zeitsch and q. Chem., 1906, 49, 341.

4 Willim, Ann. Chim. Phys., 1805, [iv.], 5, 5, Nickles, Compt. rend., 1864, 58, 537;

5 Fharm. Chim., 1864, [iv.], 1, 22, 1866, 4, 127, Ramuelsberg, Ber., 1870, 3, 360, Pogg., Annalen, 1872, 146, 592; Neumann, Annalen, 1888, 244, 329, Godeffroy, Zeitsch Ocsterr., 1806, No. 9, Pratt. Amer. J. Sei, 1895, [iii.], 49, 398, 402; Zeitsch anorg., 3 Chem., 1895, 9, 19; Johnson, J. Chem. Soc., 1878, 33, 183; R. J. Meyer, Zeitsch. anorg., 3 (Chem., 1900, 24, 321, Wallaco, Zeitsch Kryst. Min., 1911, 49, 417.

#### THAILIUM.

| Type TIX, 2MX, xH,O.                        | Type TIX, 3MY.xH3O             |
|---------------------------------------------|--------------------------------|
| TICL, 2KCl. 2H, O                           | TICl <sub>3</sub> .3NaCl,12H,O |
| TICI, 2RbCl                                 | TICI₃ั 3LaCI 8H₄Oั             |
| TICL, 2RbCl.H <sub>3</sub> O                | TICL, 3NH, CL                  |
| TICl <sub>3</sub> 2CsCl                     | TICE, 3NH,CL2H,O               |
| TICl, 2CsCl.H,O                             | TICI ( 3KCI 2H ,O T            |
| TlCl <sub>s</sub> , 2CsCl 2H <sub>2</sub> O | TICL 3RbCl                     |
| -                                           | TICI ( 3RbCl H ,O              |
|                                             | TICI] 3RbCl 211,0              |
|                                             | TICL SCSCI H <sub>2</sub> O    |
|                                             | TBr , 3RbBr 2H.O               |

The salts of the type TIN, MN  $vH_2O$  appear to possess cubic symmetry, since Pratt and Wallace have shown that the asterisked salts crystallise in combinations of the cube and regular octahedron. Rummelsherg described the potassium salts of the type 2TIX, 3MX (H<sub>2</sub>O as cubic, but according to Wallace the brounde is tetragonal (a=c=1-0.7556) and the casium chloride is described by Pratt as hexagonal (a=c=1-0.8257)

Two salts of the type TIX, 2MX H<sub>2</sub>O cry-tallise in rhombic bipyramids, isomorphous with the corresponding indium salts and with aimmonium and potassium ferric chlorides, FeCl, 2NH,ClH,O and FrCl, 2KClH,O. The data given below are due to Wallace (compare data given on p. 157)

```
TICl_2RbCl.H_2O ;  
 a b c=0 9770 1 1 4388 ; density = 3 513 at 20 ° TICl_2 2CsCl H_2O ,  
 a b c=0 9690 1 3 4321 , density = 3 879 at 20 °
```

The following salts of the type TIX, 3MX 2H,O crystallise in tetragonal bipyramids isomorphous with the corresponding indium salts -1

```
TICL 3KCl 2H,O
                     a = c = 1 = 0.7941; density = 2.859 at 20°
TICL 3NH, CL2H, O , a = c = 1 = 0.8097 , density = 2.389 at 20°
TIB: ^{3}RbBr ^{2}H<sub>2</sub>O , a = c - 1 = 0.8038, density = 4.077 at ^{2}O"
```

Thallie chloride combines with the chlorides of calcium, strontium, morphism, the, copper, and memorance, forming double salts of the type 2TlCl<sub>1</sub>MCl<sub>2</sub>6H<sub>2</sub>O<sup>2</sup> Thallie beryllium chlorode, 2TlCl<sub>3</sub>BeCl<sub>3</sub> and the amnonia. addition-product of thallie cupric wolde, 2TH, Cul, 4NH, have also been a described 3

INTERMEDIATE THALLIUM HALIDES - Thallium sesqui-chloride, Tl<sub>2</sub>Cl<sub>3</sub> or 3TlCl TlCl, is readily prepared by saturating a boiling a solution of thallie chloride with thallous chloride and cooling, or by passing a chloring into water containing thallous chloride in suspension until no n.ore; white thallous chloride is left. In each case the product should be recrystally lised from boiling water, and in the latter case the too prolonged action of ... chlorine should be avoided, since it converts the spannigly soluble thallium ? sesquichloride into very soluble thallic chloride

<sup>1</sup> Wallace, loc. cet; Pratt. loc. cet., compare data given on p. 157.
2 Gewecke, Anadem. 1909, 366, 217.
3 Neumann, Annalen. 1889, 244, 329. Jorgensen, J. prakt. Chem., 1872, [ii.], 6, 82.
4 Lamy, Ann. Chem. Phys., 1863, [ii.], 67, 885; Willin, that, 1855, [iv.], 5, 5; Crookes, and Chuch. Chem. News, 1863, 8, 1; Werther, J. prakt. Chem., 180, 13, 385. [lichberling, J. admalen, 1865, 134, 11, R. J. Meyer, Zatsch. anara, Chem., 1900, 24, 321. Cushman, Marc. Chem. J., 1900, 24, 222. V. Thomas, Ann. Chem. Phys., 1907, [viii.] 11, 204.

Thallium sesquichloride crystallises in yellow, six-sided leaflets, of density 5.9. One little of water at 25' dissolves 3 43 grams of the chloride. The aqueous solution gives with alkali hydroxides a precipitate of thallic hydroxide, and the filtrate contains thallons salt in solution

Thallium sesquichloride forms mixed crystals with thallous chloride, on account of which a number of other chlorides have erroneously been described

by Werther

Thallium dichloride, TICI, or TICI TICI, is the final product of the action of dry chlorine on dry thallons chloride at the ordinary temperature, The absorption of chlorine is at first rapid, but is subsequently very slow. Thallum dichloride is a pale yellow, hygroscopic solid which may be kept indefinitely in an atmosphere of dry chlorine at the ordinary temperature without any further absorption of the gas occurring water into thalbe chloude and thallium sesquichloude 2

Thallium sesquibromide, Tl,Br, or 3TlBr TlBr, is not so readily prepared as the corresponding chlorde, since it is decomposed by water. The best method of preparation is to moister 4 or 5 grains of thallous brounde with 1 or 2 cc of water and slowly add 0.5 e.e. of bromine About 50 grams of water are then added, and the liquid brought to the boiling point. The slight residue of thallous bromde is filtered off, and the filtrate caught in a vessel that is cooled by immersion in running water Thallium sesquibrounde separates out in bright red crystals. Water de composes the bromide into thallic bromide and a solid the composition of which depends upon the relative amounts of sesquibronide and water, but which is intermediate between that of the sesquibrounde and thallous brounde. Thathum sesquibrounde forms orange yellow or orange red mixed crystals with thallous brounde 3

Thallium dibromide, TIBL, or TIBr TIBL, is the final product of the action of dry bromme on dry thallons bromide, and is best prepared by heating thallons bromide with bromine in a scaled tube at 100 for six hours, opening the tube, and removing the excess of bromme by heating. The dibromide is also produced by the action of excess of a chloroform or carbon tetrachloride solution of bromine on thallons brounde, by the prolonged action of an excess of bromme dissolved in carbon disalphide on thallous bromide, and by the decomposition of the hydrates of thallic brounde

Thallum dibromide is a dark yellow, crystalline solid readily decomposed by water 4

Thallium chlorobromides.5 — The following intermediate chlorobroundes have been described -

| Type TlX <sub>2</sub> .                         | Type Tl <sub>2</sub> X              |
|-------------------------------------------------|-------------------------------------|
| Tl <sub>s</sub> Cl <sub>s</sub> Br <sub>s</sub> | Tl,Cl,Br,                           |
| Tl Cl Br                                        | TLCLB                               |
| TIČIBr *                                        | Tl <sub>i</sub> Cl <sub>2</sub> Br, |
| Tl.Br Cl                                        |                                     |

<sup>&</sup>lt;sup>1</sup> Abegg and Spencer, Zeitsch anorg. Chem , 1905, 44, 379.

Aberg and Spencer, Zettsch enorg. Chem., 1905, 44, 379.
 Lamy, loc ett., Thomas, loc ett.
 Thomas, loc ett., Willin, loc ett.
 Willin, loc, ett., Thomas, loc ett., and Commt. rend., 1901, 132, 1487, 1902, 134, 545; Bull. Noc. chem., 1902, [m], 27, 471, 474, 481.
 R. J. Meven, loc ett., Cushman, loc ett., and Amer Chem. J., 1901, 25, 505. Thomas, Ann. Chem. Phys., 1907, [vin.], 11, 204, Compt. rend., 1900, 130, 1316, 131, 392, 1208; 1901, 132, 80, 1487; 133, 735, 1902, 134, 515, Bull. Soc. chem., 1902, [iii.], 27, 471, 474, 481.

Type TlX, -(a) The compound Tl,Cl,Br, may be prepared by treating, thallous chloride in solution with excess of bromme, and concentrating the solution of thallic chlorobramide TICIBi, over sulphinic acid in vacuo. syrupy liquid is produced which on further concentration begins to decompose, and yellow crystals of Tl<sub>i</sub>Cl<sub>i</sub>Br<sub>i</sub> are deposited. Any thallic hydroxide produced is separated by documentation while there is still a little liquid left, and the crystals are then left over sulphuric acid until they are dry. The compound TLCLB, forms sulphin yellow, orthorhombic ('prisms, and melts at 165" to a yellow liquid. It is decomposed by water with the production of impure  $\mathrm{Tl}_1\mathrm{Cl}_4\mathrm{Br}$  , which may be purified by recrystallising it from hot water

(b) The compound  ${\rm Tl}_3{\rm Cl}_4{\rm Br}_2$  is made by the method outlined for  ${\rm Tl}_3{\rm Cl}_2{\rm Br}_4$ but the starting materials are thallons bromide and chlorine. It resembles the proceding compound, and is decomposed by water with the production of ThCLBr,

(c) The compound TICIBL is obtained by heating thallous chloride with bromme

(d) The compound TLBr Cl was obtained by Cushman by the action of aqueous thallic brounds on the compound Tl<sub>1</sub>Br<sub>2</sub>Cl<sub>4</sub>. It crystallises in vallow

Type Tl<sub>2</sub>X<sub>3</sub> - 6) One method for the preparation of the compound ThCl Br, has been given under (a) above. It is also said by Cushingar to be produced by boiling a solution of thallie chloride with thallons brounde and cooling, or by boiling a solution of thallie brounde with thallous chloride. It is also produced when a limited quantity of bromme is added to thallous chloride suspended in cold water, the yellow solid produced dissolved in builing water, and the filtered solution slowly cooled. The filtrate from the pointd Tl\_Cl\_Br\_p in six sided plates, but mixed with needles of the compound Tl\_Cl\_Br\_. deposited crystals, when concentrated and cooled, deposits more of the con-

(t) One method, due to Thomas, for the production of ThCl, Br, bas just been mentioned, but it is difficult to obtain the compound free from admixed ThO Br. R. J. Mover and Cushman have also described processes for the preparation of TLCLBr, from thallons chloride, bromme, and water

(q) The preparation of TLO By from thallors brounde, chlorine, and water has been described by R. J. Meyer

Each of the compounds TI<sub>1</sub>CI<sub>2</sub>Br<sub>p</sub> TI<sub>4</sub>CI<sub>4</sub>Br<sub>p</sub> and TI<sub>4</sub>CI<sub>4</sub>Br<sub>p</sub> forms dark orange-yellow crystals, sparingly soluble in water. With rise of temperature the crystals rapidly become deep red in colon. The two in thods given by Cushman for the preparation of Tl<sub>1</sub>Cl Br<sub>4</sub> are said by that anthor to yield stereo-isomeric forms of the compound, but at present this cannot be regarded as proved

Thallium sesqui-iodide, TLL, or 3TH TH, has been described by Knosel 1 The compound Tl,1, or 5TH TH, was discovered by Strecker, and later, prepared by Jorgensen 2. It has been shown by Abegg and Martland that thallous rodide passes into this compound at 25' in the pre-ence of a solution of rodine in water or potassium rodide solution, provided that the concentration of free iodine in the liquid phase exceeds 0.0019 gram per htre, but does not exceed 0.082 gram per litre. When the concentration of free roduc exceeds the latter value, the compound TI,I, passes into thallie

# ALUMINIUM AND ITS CONGENERS.

odide. The compound Tl,I, is a black solid, sparingly soluble in water and organic liquids with partial decomposition.

Thallous chlorate, TCIO,—An aqueous solution of this salt is obtained by dissolving thallium in aqueous chloric acid, or by mixing equivalent quantities of barnin chlorate and thallous sulphate in solution. The chlorate separates from aqueous solution in microscopic prisms of specific gravity 5 047.2 The solubility, in grains of salt per 100 grams of water, is as follows:—

One litre of the saturated solution at 20' contains 38 53 grams of thallous chlorate (Noyes and Farrell) When heated to 186°, the chlorate explodes 8

Thallons chlorate is monochnic and isomorphous with potassium chlorate; and these two salts form an incomplete series of mixed crystals with one another.

Thallic chlorate,  $\mathrm{Tl}(\mathrm{ClO}_1)$ ,  $4\mathrm{H}_2\mathrm{O}$ , forms deliquescent, rhombic crystals <sup>5</sup> Thallous perchlorate,  $\mathrm{TlClO}_0$ , may be obtained in aqueous solution by dissolving thalloun in aqueous perchlorate and thallous sulphate. It crystallises in rhombic tablets (a b c=0.7978;  $\pm$ 1:12898) of density 1814, and begins to decompose at 300%. The solubility, in grains of anhydrous salt per 100 grains of water, is 10.0 at 15% and 166.6%, at 100%.

Thallous perchlorate forms a complete series of mixed crystals with potassium perchlorate, with which it is isomorphous.

Thallic perchlorate, Tl(ClO<sub>1</sub>), 6H<sub>2</sub>O, forms white, hygroscopic crystals. 

Thallous bromate, TlBrO<sub>p</sub> obtained by dissolving thallous oxide or carbonate in aqueous bromic acid, crystallises from aqueous solution in needles. It decomposes at 100° and detonates when heated to 150° At 20°, 100° graphs of water dissolve 0, 347 of the hygoria, at 40°, 0,741 graphs.

100 grams of water dissolve 0.347 of the bromate, at 40°, 0.741 gram 9

Thallic bromate, Tl(BrO<sub>3</sub>), 3H<sub>2</sub>O, forms white, hygroscopic crystals <sup>10</sup>

Thallous iodate, TlO<sub>3</sub>, is obtained as a white, crystalline precipitate

Thallous iodate, THO<sub>3</sub>, is obtained as a white, crystalline precipitate when a solution of an alkali iodate is added to a solution of a soluble thallous salt. It begins to decompose above 150°, with the evolution of oxygen and iodine. Ditte has described a hemily drate, 2THO<sub>1</sub> H<sub>2</sub>O <sup>12</sup>

A saturated solution at 20° contains 0 058 gram of thallous indate per 100 grams of water. 13

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1 Abegg and Matland, Zeitsch anorg (Them., 1905, 49, 341.
2 Crookes, Chem News, 1863, 8, 159, J Muir, Trans (Them Soc., 1876, i 857.
3 Gowocke, Zeitsch anorg Chem., 1912, 75, 272.
4 Stortenboker, Rec. trar chim., 1905, 24, 53; Roozeboom, Zeitsch, physikal Chem., 1891, 8, 513.
3 Gewecke, loc. cit.
6 Rooce, J. Chem. Soc., 1866, 19, 504, cf. Carnelley and O'Shea, Trans. Chem Soc., 1884, 45, 409.
7 Stortenboker, loc. cit.
3 Gewecke, loc. cit.
3 Gewecke, loc. cit.
4 Gowecke, loc. cit.
4 Gowecke, loc. cit.
5 10 Gewecke, loc. cit.
5 11 Gewecke, loc. cit.
5 12 Gowecke, loc. cit.
6 13 Gowecke, loc. cit.
7 13 Gowecke, loc. cit.
7 14 Gowecke, loc. cit.
7 15 Gowecke, loc. cit.
7 16 Gowecke, loc. cit.
7 17 Gowecke, loc. cit.
8 18 Ditter, Ann. Chim. Phys., 1890, [vi.], 21, 145.
18 Ditter, Ann. Chim. Phys., 1890, [vi.], 21, 145.
18 Ditter, Ann. Chim. Phys., 1890, [vi.], 21, 145.
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Thallic iodate, Tl(IO<sub>3</sub>)<sub>3</sub> — Soluble hydrates of this salt, containing 2H and 12H,O, have been prepared !

Thallous permanganate, TIMnO, prepared by double decomposition between barum permanganate and thallous sulphate, crystallises from aqueous. solution in large, black prisms 2

# THALLIUM AND THE OXYGEN GROUP.

Oxides and Hydroxides of Thallium. Two basic oxides of thallium are known, thallows oxude, TI,O, and thallie oxude, TI,O, In addition, thallothallie or de, Tl<sub>2</sub>O Tl<sub>2</sub>O<sub>3</sub> or TlO, is known, and an oxide of the formula, Tl<sub>3</sub>O<sub>5</sub> has been described.

Normal thallows hydrorade, TIOH or TI,O H,O, and monohydrate of thallie or de or metathallic hydrocide, TIO OH or TI,O, H,O, are also known.

Thallous oxide, TLO, is produced when thallium oxidises in air or oxygen at comparatively low temperatures. In warm air the oxidation of the metal proceeds rapidly, and at temperatures approaching dull redness thallie oxide commences to be formed in appreciable amount. Thallous oxide is best prepared by dehydrating thallons hydroxide at 100°. It is a heavy, black powder, which rapidly absorbs water, producing thallous hydroxide, and reacts with acids to produce thallous salts. It is attacked by chlorue and bromme with the production of chlorides and broundes of thallium and free oxygen At about 300 it melts to a higher hapidly attacks glass When heated to redness in hydrogen, thallons oxide is slowly reduced to metal, the reduction proceeds more readily in a stream of carbon

Thallous hydroxide, TIOH, cryst illises from aqueous solution in yellow . needles. A solution of the hydroxide is readily prepared by shaking an excess of thallium timings with water in a vessel through which a stream of oxygen is passing, or by double decomposition between thallous sulphate

Thallous hydroxide is readily dissociated into thallous oxide and water ;-2TIOH TIO + H.O.

the equilibrium pressure reaching the value 760 mm, at 139°, according to the measurements of Bahi. The hydroxide also darkens in colour in vacuo. when exposed to light, but the nature of the chemical change is unknown.

Thallous hydroxide is readily soluble in water and alcohol One litre of the saturated aqueous solution at t' contains  $\epsilon$  grams of the hydroxide, where x and t are as follows = 3

0° 20" 30' t° 10° 501 60° 70° 80° x 254 4 352 7 402 8 495 0 594 5 733 1 888 8 1062 6 1260 8 1505 0

Thallous hydroxiders a comparatively strong base ' The aqueous solution' is colourless and has a strongly alkaline reaction. It precipitates metallio

<sup>1</sup> Gewecks, loc. cil
2 R. J. Meyer and Best, Zeitsch, anorg. Chem., 1899, 22, 169
2 Lamy, Ann. Chim. Phys., 1864, [m. ], 67, 385. Flemming, Jahresher., 1808, p. 250;
3 Lamy, Eds., 1890, 23, 788. Ranter, Annalen, 1892, 270, 249.
4 Bahr, Zeitsch, anorg. Chem., 1911, 71, 79
5 Calculated from the data given by Bahr, loc cit.
6 Ostwald, J. prakt. Chem., 1886, [m.], 33, 352.

hydroxide by double decomposition, readily absorbs carbon dioxide, and attacks glass and porcelain. Accordingly, unless special care is taken in its preparation, thallous hydroxide is always contaminated with a little carbonate.

Thallous hydroxide is oxidised to dark brown-thallie hydroxide by ozone. The reaction has been proposed as a test for ozone, but is given by other oxidising agents, eg hydrogen peroxide  $^4$ 

The hydrate, TiOH H<sub>2</sub>O, has been obtained, but the conditions under which it is produced are not known.

Thallic oxide or thallium sesqui-oxide, Tl<sub>2</sub>O<sub>p</sub> is produced by the action of an or oxygen upon thallium at temperatures between a dull red heat and 700°, and may be prepared by dehydrating thallic hydroxide. It is obtained in the crystalline form by fusing thallous sulphate, intrace, or chromate with excess of potassium hydroxide at a dull red heat and extracting the mass with water, or by heating thallous intrate to 450.6°. The crystals have a density of 9.97

When hydrogen perovide is added to a cold solution of a thallous salt containing excess of alkali hydrovide, brown thallie oxide is precipitated, which soon becomes crystalline and has a density of 9.65 at 21°. When precipitation is effected at 80, the oxide obtained is black and amorphous, and has a density of 10.19 at 22°. At temperatures above 100°, the brown variety becomes black, each form contains a little water which is not completely eliminated below 500°.

Thallie oxide is deposited upon the anode when a neutral solution of thallons sulphate or intrate is electrolysed between platinum electrodes and the anodic potential difference (between electrode and electrolyte) exceeds 143 volts. When the potential difference at the anode exceeds 181 volts for thallons intrate or 2.27 volts for thallons sulphate, thallie hydroxide is deposited 8.

Thallic oxide melts at  $725^{\circ}\pm10^{\circ}$ , begins to decompose into thallous oxide and oxygen at 800°, and decomposes rapidly at  $1000^{\circ}$ °. It is reduced by hydrogen and by carbon monoxide at a red heat  $^{10}$ °. It is insoluble in water and in alkalies, it reacts with hydrochloric acid, producing thallic and thallous chlorides and chlorine , it is scarcely affected by cold sulphiric acid, and is attacked by the hot acid with the production of thallous sulphate and oxygen  $^{11}$ 

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1 Lamy, loc. ett., Canelley and Walker, Trans. Chem. Soc. 1888, 53, 59, 28 Schonhem, J. peak! Chem., 1867, 101, 321, Lamv, Bull. Nov. chem., 1869, [n.], 17, 210.

3 Schone, Annalen. 1878, 196, 58.

4 Willin, Ann. Chem. Phys., 1865, [n.], 5, 5.

5 Lepheric and Luchand, Compit. rend., 1891, 113, 196.

6 V. Thomas, thal., 1904, 138, 1697

7 Rabe, Zettsch. annay. Chem., 1906, 48, 427, 1906, 50, 158, 1907, 55, 130.

8 M. Bose, Zettsch. annay. Chem., 1905, 44, 237, see also Crookes, Proc. Roy. Noc., 1862, 12, 159; Bottger, Annalen, 1863, 127, 175, Wohler, ind., 1868, 146, 233, 375. Fl. manag., Iso; Bottger, Annalen, 1863, 127, 175, Wohler, ind., 1868, 146, 233, 375. Fl. manag., Elektrochem., 1903, 9, 523; Lorenz, Zettsch. annay. Chem., 1896, 12, 439. According to Gallo and Cenn. (Alte. R. Accad. Lines, 1908, [v.], 17, n. 276), the oxide deposited on the sanode when thallous sulphate solution containing a trace of sulphure acid is electrolysed has, when dired at 160° to 200°, a composition corresponding to the formula Th.O.

8 Rabe, loc. ett.; cf. Carnelley and O'Shea, Trans. Chem. Soc., 1581, 45, 409.

9 Worther, J. prakt. Chem., 1864, 91, 385. Flemming, loc. ett.; see also Winkler., 1890, 23, 788, Bottger, Dind. polis. J., 1870, 197, 379.

Lamy, Ann. Chem. Phys., 1863, [iii.], 67, 335.
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Thallic hydroxide, 1 TIO.OH or TI2O, H2O, is obtained by precipitating? a solution of a thallic salt with a slight excess of alkali hydroxide or ammonia washing the precipitate, and drying it at the ordinary temperature in the air.2 It is also produced by the action of alkali hypochlorite on an alkaline solution of a thallons salt  $^3$ 

Thallic hydroxide is a reddish brown powder. When heated to 110" 120" it is completely delivdrated to thallie ovide, and it is also delivdrated when heated in contact with the liquid from which it has been precipitated 5. The freshly precipitated hydroxide reacts readily with acids, but, after it has been dried, solution in acids is slow, and is accompanied by partial reduction. The hydroxide is practically insoluble in water, its solubility product 6 at 25 being only  $10^{-51}$ 

Thallothallic oxide, TIO or TI O TLO, is obtained as a blinsh black, lustrous precipitate when 3 per cent. hydrogen perovide is added to a solution of thallous sulphate which has been made strongly alkaline and cooled to

It should be wished with cold alcohol and then with other oxide absorbs earbon dioxide, torning thallie oxide and thallons carbonate, it absorbs ovegen at the ordinary temperature, forming thallie oxide, and it is decomposed by cold water, producing thallous hydroxide and thallie oxide?

Sulphides of Thallium When mixtures of thallium and sulphur arg fused, those rich in thallium separate into two non-miscible layers of thallous. sulphide, TLS, and thallow, while those rich in sulphin form two layers consisting of thallium pentasulphide and sulphin Mixtures containing more sulphur than is required for thallous sulphide and less than is required for the pentasulphide form only one liquid phase, and solid solutions appear to separate when they are cooled. The nature of these solid solutions is not known possibly a thallothallic sulphide, TLS TLS, also exists, but has no melting point S

Thallous sulphide, TLS, is obtained as a black precipitate by passing hydrogen sulphide into, or adding ammonium sulphide to, a solution of a thallous salt. In the presence of a trace of sulphurie acid, hydrogen sulphide precipitates the sulphide from cold solutions as uncroscopic tetrahedra." amorphous sulphide becomes crystalline when heated for some hours to  $150^{\circ}/200^{\circ}$  with excess of colourless ammonium salphide  $^{10}$ 

Thallous sulphide forms bluish black crystals of density 8.0. It melts at 448°, and is very brittle. Heated in hydrogen, it is reduced to thallium. It

- --- --

<sup>&</sup>lt;sup>1</sup> A hydroxide of the formula ThOH), stable at 310°, has been described by Catargie (Chem. News, 1889, 60, 113°, but the formula given is based solely on an estimation of the thallium present in the substance. <sup>2</sup> Lamy, Ame Chem. 1863, [m.], 67, 785°, Crookes, J. Chem. Soc., 1864, 17, 112; Werther, J. prakt. Chem., 1864, 91, 385°, Carnelley and Walker, France Chem. Soc., 1888, 288.

<sup>53, 88,

3</sup> Steecker, Annalen, 1865, 135, 207, Willin Anna Chim Phys., 1865, [iv.], 5, 5; see
also Schonbern, A peakt Chem., 1861, 93, 35, Weltzee, Annalen, 1866, 138, 129

4 In heating thalbe oxide on bedroude, care must be taken to prevent the access of gaseous products of combustion to the substance

gaseous products of combustion to the substance

R. J. Meyer, Zeitsch anory Chem., 1900, 24, 321

Abegg and Spencer, Zeitsch anory Chem., 1995, 44, 379

Rabe, Zeitsch anory Chem., 1908, 58–23.

Pilabon, Compt. rend., 1907, 184, 118; Ann. Chem. Phys., 1207, [viii., 17, 526]

Pilabon suggests the interim date sulphide Thes.,

Phebberling, Annalen, 1865, 134, 11

Stanck, Zeitsch anory Chem., 1898, 17, 117.

is readily decomposed by dilute mineral acids, but not by acetic acid, and the moist sulphide rapidly oxidises in the air. It is insoluble in animonium sulphide, alkalı hydrovides, carbonates, and cyanides.

Thallous sulphide has been obtained in colloidal solution. 1

Thallous sulphide forms definite compounds of the formulæ 3Tl2S.As2S5 (or Tl<sub>3</sub>AsS<sub>1</sub>) and 2Tl<sub>2</sub>S.SnS<sub>2</sub> (or Tl<sub>3</sub>SnS<sub>1</sub>) with assence and stannic sulphides. They may be employed for the gravinetic estimation of thallium. With reservous, antimonious, antimonic, and stannous sulphides, thallous sulphide forms series of solid solutions. They are precipitated from mixed thallous and arsenic, antimony and tin solutions by alkali sulphides.<sup>2</sup> Thallous sulphide also forms a double sulphide of the formula TLS 4CuS with cupric sulphide, and two series of solid solutions probably of the compound PLS. 2CuS in TLS 4CuS and TLS in TLS 2CuS.<sup>3</sup>

Thallic sulphide, TLS, is not precipitated from a thallic salt by sydrogen sulphide, thallous sulphide and sulphur being produced <sup>1</sup> It may,

towever, be prepared by fusing thallium with excess of sulphur and removing excess of the latter by distillation 5. It is a black solid, hard and brittle below 12°, soft and plastic above that temperature. It dissolves in warm dilute sulphuric acid with the evolution of hydrogen sulphide, no sulphur separating us such.

With thallows sulphide, thallie sulphide forms a series of solid solutions.6 The compound potasseum thalleum sulphide, Tl28, K28, is obtained by fusing cogother thallous sulphate (1 pt.) with sulphur (6 pts.) and potassium parbonate (6 pts.), and washing the fused mass with water. It is a reddishrown, crystalline solid, insoluble in water.7

Thallium pentasulphide, Tl.S., is said by Pélabon to melt at 127°, out, if quickly cooled, to remain pasty at ordinary temperatures. When neated above the melting-point it readily loses sulphir. A crystalline pentasulphide is said to be formed by digesting thallous chloride with concentrated ammonium polysulphide solution, 8 but, according to Hawley, there is no evidence that the substance thus obtained is a definite pentasulphide, but there exists a series of solid solutions containing thallium and sulphur, in which there is more sulphur than is required for a sesquisulphide.

Thallium selenides —According to Mirakann, thallium forms thallous selenide, Tl.Se, melting at 368°; thallothallic selenide, Tl.Se Tl.Se g. melting at 310°, and thallium sesquiselenide, Tl.Se, decomposing at 265°. The equilibrium diagram is complex, there being two gaps of miscibility in the liquid state. The selenides are brittle solids. Thallous

Minssinger, Bull Noc. chim., 1888, [n], 49, 452
 Hawley, J. Physical Chem., 1906, 10, 654. J. Amer. Chem. Soc., 1907, 29, 1011., cf.
 Bruneing, Chem. News, 1868, 17, 138
 Bruner and Zawadski, Bull. Acad. Sec. Cracow, 1909, p. 312.

Cf. Strecker, Annalen, 1865, 135, 207
Carstanjen, J. prakt. Chem., 1867, 102, 65, 129
Hawley, loc cit, of Schnoider, vide infra, Carstanjen, J. prakt Chem., 1867, 102,

nawiey, or cit, cf Schneider, vide inita, Carstanjen, J. prakt Chem., 1867, 102, 15, 159.

7 Schneider, J prakt. Chem., 1870, [n], 2, 164; 1874, [n], 9, 209, 10, 55, 1890, [n], 2, 805; Pogy. Annalen, 1870, 139, 661; 1875, 153, 588; Kruss and Soloreder, Eer, 1886, 159, 2786.

8 K. A. Hofmann and Hochtlen, Eer., 1903, 36, 3090.

9 Murakami, Mem. Coll Sci. Tokno, 1915, 1, 153. of Pélsbon, Compt rend, 1907, 145, 186; Ann Chim. Phys., 1909, [vul], 17, 526; Carstanjen, loc cit., Fable, Ann Chim. Phys., 1887, [vi.], 10, 538, Kuhlmann, Bull. Soc. chim., 1864, [n], 1, 380

selenide may be precipitated by passing hydrogen selenide into aqueous thallous carbonate. It is decomposed by acids

Thallium tellurides - The system thallium - tellurium has been studied by the thermal method by Pélabon and by Clukashigé. Thallous telluride, Tl,Te, melts at \$12°, and is almost non miscible with thallium. The telluride Tl,Te<sub>2</sub> (or Tl,Te<sub>3</sub>!) melts at \$12° Thallothallic telluride, Tl,Te<sub>1</sub>Tl<sub>2</sub>Te<sub>2</sub> which crystallises in long needles, is not stable in contact with a liquid phase of the same composition

Thallous sulphite, TLSO, may be prepared by adding sulphurous acid to thallous hydroxide or by mixing solutions of thallous sulphate and sodium sulphite, when thallous sulphite is precipitated. It forms white crystals of density 6427 at 20°, and is insoluble in alcohol. It can be recrystallised from warm water, 100 parts of water at 15° dissolve 3.34 of

Thallous thiosulphate, TLS,O., is obtained as a white crystalline precipitate by adding potassium thiosulphate to a solution of a thallous salt—It can be crystallised from hot water, and begins to decompose at 130°—Sodium thallows thresulphate, 2TLS,0, 3Na<sub>2</sub>S,0, 8H<sub>2</sub>O, is known <sup>3</sup>

Thallous sulphate, TLSO, is prepared by dissolving thalloum in hot sulphuric acid, or by adding thallous hydroxide or carbonate to aqueous sulphuric acid, evaporating, and crystallising

Thallous sulphate crystallises in Thombic prisms (a b c=0.5555 1: 0.7328) of density 6.765 at 20° 4. It is therefore isomorphous with potassium sulphate (for which a  $b \cdot c = 0.5727 \cdot 1 \cdot 0.7418$ ), and these two salts form a continuous series of mixed crystals  $^6$ . Thallous sulphate melts at  $632^{\circ}$ ,  $^7$ and at temperatures above dull redness it volatilises. The mean refractive index, 18708 for the D line, is extremely high, the value for potassium# sulphate being 1:1952 (Tutton)

One hundred grams of water dissolve the following amounts of thallous sulphate -- "

At a red heat hydrogen reduces thallous sulphate to a mixture of sulphide and metal, and ammonia reduces it to thallous sulphide."

Numerous double sulphates containing thallous sulphate are known, With the sulphates of the tervalent metals aluminium, chromium, iron, vanadium, gallium, and shodium, thallous sulphate forms alums, of the

.... ... ... ...

Pélabon, los cit., Chika-lugé, Zeiteh. ann q. Chem., 1912, 78, 68., Fabre, Ann. Chim. Phys., 1888.
 I. 14, 115. According to Chika-lugé, thallous telluride does not exist, but this conclusion is questionable
 Scubert and Elten, Zeitsch annry. Chem., 1892, 2, 434, 1893, 4, 44; Rohrig, J. makt., Chem., 1888, [u.], 37, 217
 Ilebberling, Annalen, 1865, 134, 11; Werthu I. prakt. Chem., 1861, 92, 130., Vortemann and Padberg, Ber., 1869, 22, 2637. Joshum, Chem. Zent., 1885, p. 612
 Tutton, Proc. Roy. Soc., 1907, A., 79, 351.
 Von Lang, Phil. May., 1863, [u.], 25, 248., Annalen 1863, 128, 76., Tutton, loc. est.
 Stortenbeker, Ec. trav. ch.m., 1905, 24, 53
 Cainelley and Williams, Trons. Chem. Soc., 1878, 33, 281
 Earl of Berkeley, Phil. Trans., 1904, A., 203, 211., Sudell, Solubilities (Crosby, Lockwood & Son, 1907), p. 338, see also Noyes, Fairell, and Stewart, J. Amer. Chem. Soc., 1813, 33, 1650.

<sup>1911, 33, 1650.</sup>By Hodgkinson and French, Chem. News, 1893, 66, 223.

general formula Tl<sub>2</sub>SO<sub>1</sub> R<sup>m</sup><sub>2</sub> (SO<sub>4</sub>)<sub>3</sub> 24H<sub>2</sub>O <sup>1</sup> With the sulphates of magnesium, zine, iron, nickel, cobalt, copper, manganese, and cadmium, thallons sulphate forms a series of isomorphous, monochine, double sulphates of the general formula TI SO, MSO, 6H,O, isomorphous with the corresponding series of salts containing ammonium, potassium, rubidium, and caesium in place of thallium.2 The crystallographic constants, determined by Werther and by Tutton (zinc salt), are as follows -

| Salt                       | a .    | b | c      | В         |
|----------------------------|--------|---|--------|-----------|
| TUSO, MgSO, 611,0          | 0.7422 | 1 | 0.5002 | 106 21'   |
| TLSO, ZuSO, 6H,O           | 07113  | 1 | 0.5010 | 106 16'   |
| TĘSO, FeSO, 6H,O           | 0.7366 | 1 | 0 4964 | 105 - 52' |
| ${ m TI, SO, NiSO, 6H, 0}$ | 0.7431 | 1 | 0 4988 | 106 39'   |
| [K,SO, ZnSO, 6H,O          | 07113  | 1 | 0.5044 | 104 487   |

The density of the thalloun one salt is 3.720 at 20°, the mean refractive index for the D line is 1 6064, while that of the potassium zinc salt is 1 1859

Thallous hydrogen sulphate, THISO4, is obtained when thallous sulphate is dissolved in sulphuric acid and brought to constant weight at  $220^\circ$  to  $210^{\circ 3}$ . It may be also crystallised from a solution of thallous sulphate in moderately concentrated sulphuric acid, and exists in a monoclinic and an orthorhombic (!) form, while another modification would appear to exist, since the salt is isotimorphous with potassium hydrogen sulphate 3

The acid sulphate,  $\mathrm{Tl}_1\mathrm{H}(\mathrm{SO}_1)_{\mathfrak{D}}$  crystallises when a solution of thallous sulphate with ten equivalents of sulphuric acid is concentrated. The crystals are trigonal (a. c = 1/3.717). Another modification probably exists, since the salt is isodimorphous with the corresponding potassium salt "

Thallous pyrosulphate, TLS,O5, may be prepared by heating thallous sulphate with sulphur trioxide?

Thallic sulphate, TL(SO<sub>4</sub>), Anhydrous thallic sulphate is said by Meyer and Goldschmidt to be obtained by heating acid thalhe sulphate to 220°. The heptahydrate, TL(SO<sub>1</sub>), 7H<sub>2</sub>O, has been described by Strecker, 8 but later experimenters have failed to prepare it

Thallie sulphate is hydrolysed by water with great readiness, brown thallie hydroxide separating out. From a hot solution of thallie hydroxide in moderately concentrated sulphuric acid a basic thallie sulphate, Th(OH)SO, 2H,O, crystallises out, accompanied by the amorphous basic salt Th(OH)SO,  $11_2$ O  $^{10}$ 

See pp. 81, 82, 83, 148, 160, the subsequent volumes of this series; and Lainy, Ann-Chim Phys., 1863, (iii.), 67, 385, Soiet, Aich Sei phus ant., 1881, [iii.], 12, 553, 1888, [iii.], 20, 520; Soiet and Duplic, thid., 1889, [iii.], 21, 90; Spring, Ber., 1881, 16, 2723; Polis, Ber., 1880, 13, 397, Cosa, Nuovo Cimento, 1870, [ii.], 3, 75, Ber., 1878, 11, 811; Stolla, J. Chem Soi., 1874, 27, 873, Locke, Amer Chem J., 1901, 26, 166.
 Weither, J. piakt Chem., 1864, 92, 128, 351; Willin, Ann. Chim. Phys., 1865, [iv.], 5, 5; Lamy and Descloremy, thid., 1869, [iv.], 17, 310. Tutton, Proc. Roy. Soc., 1910,

- Browning, Amer. J. See, 1900, [iv.], 9, 137; Zeitsch. amorg. Chem., 1900, 23, 155
   Stortenbeker, Rec. trac. chem., 1902, 21, 87, 1905, 24, 53; 1907, 26, 248
   Gossner, Zeitsch. Kryst. Min., 1903, 38, 158.

- \*\*Grossner, Zettsen, Arjan, 1993, 36, 13c.

  \*\*Strotenbeker, Io. et.

  \*\*R. Weber, Rev., 1884, 17, 2502, 2707.

  \*\*Streeker, Arnalen, 1865, 135, 207

  \*\*Stereker, Arnalen, 1865, 135, 207

  \*\*Steeker, Arnalen, 1865, 128, 29, 22, 596.

  \*\*Willin, Aun. Chim. Phys., 1865, [1v.], 5, 5, Marshall, loc. etc.

Acid thallic sulphate. HgSO4 Tl2(SO4), 8H2O, crystallises out, according to Meyer and Goldschmidt, from a solution of thallic hydroxide in. concentrated sulphuric acid.

The following double sulphates containing thallie sulphate are known:— $^1$ 

Further, Marshall has prepared the basic salt of the formula Tl(OH)SO, K-SO, Thallie ammonium sulphati, TL(804) (NH4) 8048H40, crystallises in mono clinic prisms (a=b, v =0.9559–1=0.6836 ,  $\beta$  =125–38 )  $^2$ 

No alums of thellic sulphate have vet been prepared, but Piccini and Forting have prepared mixed crystals of ammonium alum and ammonium thallie alum  $^3$ 

A number of thallothallic sulphates have been described simplest compound, TLSO<sub>4</sub> TL(SO<sub>4</sub>), separates from a sulphinic acid solution of the sulphates, inived in the ratio of their molecular weights, in crystals which are said by Maishall to be probably orthorhombic. Maishall has also prepared double sulphates of the formula  $571.80_111.80_1$ , and 2TI SO<sub>1</sub> TL(SO<sub>1</sub>), while Willin has described a compound of the formula 2Tl<sub>2</sub>SO<sub>1</sub> 3Tl<sub>2</sub>(SO<sub>4</sub>)<sub>1</sub> 22H<sub>2</sub>O<sub>3</sub>

By oxidising a solution of thallous sulphate with bromine and crystallising, Meyer and Goldschmidt have prepared thallothallie bromesulphate, TlaBr SO,

Thallous persulphate. The O., prepared by double decomposition between ammonium persulphate and thallous sulphate, or by the electrolysis of thallous sulphate in a divided cell, forms monochine crystals isomorphous with the ammonium salt " It is isomeric with thallothallic sulphide, Thso, Thso,

Thallous dithionate, TLS,O6, is prepared in aqueous solution from barum dithionate und thallons sulphate in equivalent proportions. It crystallises in shining, monochine prisms (a b c 0.9292 1.0.3986;  $\beta$ = 96' 58') of density 5.573, and one grain of the salt at 18.5' dissolves in 2.39grams of water ?

Potassium dithionate forms trigonal crystals, but with thallons dithionate 4 it gives two series of mixed crystals, so that each salt is probably dimorphous. Thallons dithionate is isomorphous and forms monochine mixed

Cr. Lepsius, Chem. Zentr., 1891, i. 691 C. Lepsins, Chem. Zeatr., 1891. 1, 691
 Willin, Ann. Chim. Phys., 1865. [w.], 5, 5. Abegg, Asorquatischen Chemia (Leipzig., 1906), vol. in., pt. 1, p. 146.
 Foster and E. Smith, J. Amer. Chem. Soc., 1899, 21, 934. Marshall, chid., 1909, 22, 48. 7
 Kluss, Annalen, 1888, 246, 179, 284. Fock. Zeitsch. Keyst. Min., 1882, 6, 169.
 Stortenbeker, loc. cd.

crystals with barron dithionate. It also forms a trichine, double salt with thallous sulphate, 3Tl<sub>2</sub>S<sub>1</sub>O<sub>2</sub> Tl<sub>2</sub>SO<sub>3</sub>, and a monoclinic compound with thallous hydroxide, Tl<sub>2</sub>S<sub>2</sub>O<sub>3</sub> TlOH H<sub>2</sub>O<sub>2</sub>

Thallous selenite, Tl<sub>2</sub>SeO<sub>3</sub>, prepared from thallous carbonate and selenious acid, forms micarcous crystals, soluble in water but misoluble in alcohol or other. Acid thallous selenite, TlllSeO<sub>3</sub>, is more soluble in water than the normal salt <sup>3</sup>

Thallic selenite  ${\rm Tl}_g({\rm SeO}_g)_p$  may be obtained in white crystals by the action of selemons and on thallic hydroxide  $^4$ 

Thallous selenate, Tl<sub>o</sub>SeO<sub>p</sub> may be prepared by neutralising selenic acid with thallous hydroxide or carbonate. It crystallises from water, in which it is not very soluble, in orthorhombic prisms  $(a \cdot b \cdot c = 0.5551 \cdot 1.0.7243)$  of density 6.875 at 20°, and is therefore isomorphous with potassium selenato  $(a \cdot b \cdot c = 0.5731 \cdot 1.0.7319)$ . The mean refractive index, 1.9575 for the D-line, is very high, the value for potassium selenato being 1.5396. The salt is insoluble in alcohol and ether, it melts above  $400^{\circ}$ .6

Thallous aluminium selenium alum, Tl<sub>2</sub>SeO<sub>4</sub> Al<sub>2</sub>(SeO<sub>4</sub>), 24H<sub>2</sub>O, is described on p. 87

Of the double selenates that thallous selenate may be anticipated to form with the selenates of the bivalent metals, only the zine salt, thallous zine selenate, TI SeO, ZISCO, 6H,O, has been described. It forms monoclinic crystals (a h r=0.7479 l 0.5022,  $\beta=105^{\circ}$  54') isomorphous with the corresponding double sulphates and with potassium zine selenate (a h c=0.7458 l 0.5073,  $\beta=104^{\circ}$  12').

Thallic selenate,  $\Pi_2(SeO_1)_p$  has not been described, but ammonument and potassium-thallic selenates,  $(NH_1)_sSeO_1\Pi_2(SeO_1)_sSH_2O$  and  $K_2SeO_1$ ,  $\Pi_2(SeO_1)_pSH_2O$ , analogous to the double sulphates, are known  $^8$ 

Thallous tellurate, Tl\_TeO, obtained from thallons oxide and telluric acid, is a white, amorphous, insoluble powder 9

Thallous chromate, Tl<sub>2</sub>CiO<sub>D</sub> is precipitated when a solution of a thallous salt is mixed with potassium chromate. It is also produced by adding thallous hydroxide or carbonate to aqueous chromic acid, or by the action of ammonia on the dichromate. Thallous chromate is a yellow crystalline powder, very sparingly soluble in water, 100 grains of which dissolve 0.03 grain of the salt at 60° and 0.2 grain at 100°. In hot, concentrated potassium hydroxide solution thallous chromate is fairly soluble, and it separates out on cooling in crystals which are apparently isomorphons with potassium chromate. At a dull red heat the chromate melts, and at higher temperatures it loses oxygen.

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1 Kluss, Voc. ett. Fock, Zeitsch Kryst Min., 1888, 14, 340., Stortenbeker, loc. ett.
2 Wyrouboll, Bull Soc. chim., Inn., 1881, 7, 139, see also ibid., 1882, 5, 32.
3 Kuhlmann, Bull. Soc. chim., 1811, 11, 1, 130.
4 Manno, Zeitsch anorg Chem., 1909, 62, 173.
5 Tutton, Proc. Roy. Soc., 1907, A, 79, 351., Kuhlmann, loc. ett.
6 Gausser, Zeitsch anorg. Chem., 1910, 66, 137.
7 Tutton, Proc. Roy. Soc., 1910, A, 83, 211.
8 Fortin, E'Dross, 1902, 25, 397
9 F. W. Clarke, Amer. J. Soc., 1878, [in.], 6, 201, Dennis, Doan, and Gill J. Amer.
C'hem. Soc., 1896, 18, 970.
10 Browning and Hutchius, Amer. J. Soc., 1899, [iv.], 8, 460, Zeitsch anorg Chem.,
1900, 22, 380.
11 Lepierre and Lachaud, Compt. rend., 1891, 113, 196.
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#### THALLIUM.

Thallous dichromate,  ${\rm TL}(C_1O_2)$ , is precipitated as an orange-red, crystalline powder when solutions of a soluble thallous salt and potassium dichromate are mixed (Crookes), and is also obtained by the action of sulphuric acid on the chromate (Willin)

Thallous trichromate, Tl<sub>2</sub>Cr<sub>4</sub>O<sub>100</sub> a red, crystalline, sparingly soluble salt, is obtained by the action of intin acid on the dichromate, or of concentrated sulphuric acid on the chromate (Crookes, Willin)

Thallous chlorochromate. Cit (OTI)Cl, obtained from thallous chloride and chromic acid, forms small, prismatic crystals, decomposed by water !

Thallic chromate, Tl<sub>2</sub>(CrO<sub>4</sub>)<sub>by</sub> has not yet been prepared, but potassium thallie chromate,  $K_1C(O_1|T)_2(C(O_4))_1(H)_2O_5$  is known as a yellow, crystalline salt, decomposed by water  $^2$ 

Thallous molybdate, Tl MoO, is obtained as a white, sparingly soluble, crystalline solid by double decomposition, or by beiting a solution of a thallous solt with molybehe and. More complex molybehees of the formulæ Tl<sub>2</sub>O 11MoO<sub>9</sub> 3Tl<sub>2</sub>O 8MoO , Tl<sub>2</sub>O 4MoO H<sub>2</sub>O, and 5H<sub>2</sub>O 12MoO<sub>80</sub> have also been described

Thallous phosphomolybdate, 3TI O P<sub>2</sub>O<sub>2</sub> 20MoO , has been described by Debray , 1 and thallous silicomolybdate, 2TL0 8iO, 13MoO, rH<sub>2</sub>O, by Parametric  $^{3}$ 

Thallous tungstate, TLWO, is obtained by heating aqueous thallous

Cibonate with tringstic oxide, as a sparingly soluble, white, crystalline solid for Thallous phosphotungstate, TLOPO 12WO 4H O, has been descril ed also acid thallous silicotung state, 11,0 H O(SiO, 12WO), 9H,O 8

# Thallium uranates "

## TPAIRIUM AND THE NITROGEN GLOUP

Thallous azide overhoutride, bydrassie, 'IIN', is precipitated as a white, cry tilling solid when solutions of thidlors sulphate or nitrate and sodium azide are mixed. It is sparingly soluble in cold, more readily soluble in hot water, from which it separates on cooling in tetragonal needles (a  $\approx 1/0.5881$ ) isomorphous with the potession and cubidinin salts  $^{10}$ Thollous azide slowly turns brown in sunlight. It melts at 331 in an atmosphere of carbon dioxide, and is reduced to thalliam when heated in

Thallothallic azide, TIN, TICN  $\rho_{\nu}$  separates in reflow, or thorhombic needles when thallic hydroxide is dissolved in aqueous hydrazoic acid and the solution evaporated in vacuum over sulphuric acid. It is very explosive,

13

Leparte and Luchanl, tempt rend, 1891, 113, 196
 Hawley, J. Amer. Chem. Soc., 1997. 29, 309
 Delafont one, Arch. Sec., 1997. 29, 309
 Delafont one, Arch. Sec., 1997. 29, 309
 Delafont one, Arch. Sec., 1995. 46, 428. Wenne, chal., 1912, 78, 298
 Ibelany, Compt. cond., 1868. 66, 701
 Parmentier, Compt. rend., 1881, 92, 1234
 Fleroning, loc. cit., see also Schafer, Zeitsch. amorg. Chem., 1994. 38, 142
 Pechard. Ann. Chim. Phys., 1891, [vi.], 22, 187
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 Roscolousch. Zeitsch. Krys. Men., 1996., 33, 99
 Dennes, Doan, and Gill, J. Amer. Chem. Soc., 1896, 18, 970., Curtues and Ressom, J. prakt. Chem., 1898, [n.], 58, 261.
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Hot water decomposes the compound, thallic hydroxide being precipitated and thallons azide remaining in solution 1

Thallium nitride, Ti,N, is obtained as a black precipitate when liquid ammonia solutions of thallors intrate and potassium aimde are mixed -

$$3\text{TINO}_3 + 3\text{KNH}_2$$
  $\text{TI}_3\text{N} + 3\text{KNO}_3 + 2\text{NH}_2$ 

It dissolves in a liquid ammonia solution of ammonium intrate, ie forming thallous intrate .-

$$TI_1N + 3NII_3$$
,  $NO_3 = 3TINO_3 + 4NII_3$ 

and also dissolves when excess of potassium amide is added. The latter solution when evaporated at  $-10^{\circ}$  deposits yellow crystals of potassium annule and depotassium ammonothallite, TINK<sub>2</sub> 4NH<sub>3</sub>, a compound which dissociates at 20° in vacuum, leaving the compound TINK<sub>2</sub> 2NH<sub>3</sub>  $^2$ 

Thallium phosphide .- According to Lamy, thallium and phosphorus combine when heated, but no definite compound has yet been isolated

Thallium arsenide - When thallium and arsenic in atomic proportions are heated together, a white, soft, crystalline mass is obtained, which is decomposed by cold acids, arsine being evolved  $^{1}$ 

Thallium antimonide resembles the arsende and is prepared in a sımılar manner

Thallous natrite, TINO2 is prepared by mixing solutions of thallous sulphate and barmm intrite, filtering, evaporating the filtrate in vacuum, and adding alcohol. It forms orange red crystals. The concentrated aqueous solution is yellow, but dilute solutions are nearly colourless. The dry salt has a low melting-point, it is unchanged at 140, but at higher temperatures decomposes into thallous oxide and oxides of introgen, its stability being much inferior to that of the alkali nitrites 6

Triple intrites containing thalling are known 7

Thallous nitrate, TINO, - Thallium dissolves readily in intric acid, and by dissolving the metal, thallous oxide, hydroxide, or carbonate in mirro acid, a solution of thallous intrate is obtained from which the salt crystallises in the anhydrous state in orthorhombic prisms (a b = c = 0.5109 + 1.0 = 0.6507) of density 5 55 8. At 72 8, the thombic substance changes into a thombohedral form, which at 142.5' changes to a cubic modification, and this, at 205', melts 9. On cooling, the reverse changes take place. The transition at 72 8° is a very slow process. At temperatures above 300° decomposition of the intrate takes place, it is rapid at 150', crystalline thallic oxide being left, and oxygen, introgen, and oxides of introgen being evolved in A little of the intrate volatilises unchanged

- :

<sup>1</sup> Dennis, Doan, and Gill. loc cit
2 Franklin, J. Physical Chem., 1912, 16, 682
3 Willin; Carstanjen, Flemming, Jahresber, 1868, p. 250.
4 Carstanjen, J. prakt. Chem., 1867, 102, 65, 129
5 Carstanjen, loc cit., Omodel, Jahresber, 1892, p. 153.
6 Vogel, Zeitsch. amorg. Chem., 1903, 35, 403, Thomas, Compt. rend., 1904, 138, 1697;
Ball and Abrain, Trans. Chem. Soc., 1913, 103, 2132
7 Przibylla, Zeitsch. anorg. Chem., 1898, 18, 448, Ball and Abrain, Trans. Chem. Soc., 1913, 103, 2116

Yraibylle, Zeikek anory, Chem., 1898, 18, 448., Ball and Abram, Trans Them. Soc., 1913, 103, 2116
 Lamy, Ann Chim. Phys., 1863. [in.], 67, 385.; Lamy and Descloreanx, that, 1869, [iv.], 17, 310., Miller, Proc. Roy. Soc., 1865, 14, 555.
 Van Eyk, Zeitsch. physikal. Chem., 1905, 51, 721, 1899, 30, 430; Gossner, Zeitsch. Kryst. Min., 1903, 38, 110, Retgers. Jahrh Min., 1896, Il. 183.
 Thomas, Compt. rend., 1904, 138, 1697.

Thallous nitrate is insoluble in alcohol Its solubility in water is given in the following table -1

t °C. 0° 10° 20° 30° 40° 50°  $60^{\circ}$ 80° 90 100' 105° 70` **a** 376 586 872 1251 1733 2333 3155 4101 5260 6666 8051 8559 **b** 3 91 6 22 9 35 14 3 20 9 30 4 46 2 69 5 111 0 200 0 414 0 594 0

A saturated solution in contact with excess of the salt boils at 105°.

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Mixtures of molten thallous nitrate and silver, mercurous, or mercurio intrate are useful liquids in which to effect the separation of numerals by density differences. The following melting points and densities are due to Retgers - 2

Composition of salt  $\frac{\mathrm{TINO}_4}{5.3} = \frac{\mathrm{TIAg(NO_4)_2}}{1.8} = \frac{\mathrm{TIHg(NO_4)_2}}{5.3} = \frac{\mathrm{TI_2Hg(NO_2)_4}}{5.0}$ Density of liquid Melting-point °C

When thallous intrate is dissolved in cold intra acid (density, 15) and the solution is cooled below 0° C acid thallous nitrate, TINO, 2HINO, crystallises out. The compound melts below the ordinary temperature 3

With sodium aitiate, thallous nitrate forms neither a compound nor mixed crystals 4

From a study of the crystals deposited from a solution containing both politsium nitrale and thallons intrate, it has been concluded that these two salts are isodimorphous. The fusion curve indicates that two series of solid solutions separate out The series containing excess of thallous salt is cubic, the other series being rhombohedral. These mixel crystals are transformed at lower temperatures into other crystalline modifications, and the various regions of at dulity are indicated in the diagram (fig.  $187^{\circ}$ 

Rubidium nitrate, and also cascum intrate, forms a continuous series of cubic mixed crystals with thallous intrate. The cubic crystals become thombohedral at lower temperatures, and those with a very ligh thallium content become orthorhombic at still lower tenderatures. Cubic crystals always separate from the fused maxtures, except those very rich in rubidium, which crystallise first in rhombohedial crystals different from those already mentioned?

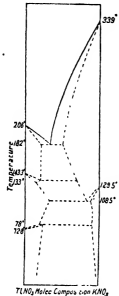
Ammonium intrate forms a continuous (t) series of cubic naved crystals with thallons intrate These crystals separate from fixed involutes of the nitrate on cooling. Two other series of mixed crystals are stable at lower temperatures 8

<sup>1</sup> Earl of Berkeley Ind Trans, 1904, A, 203, Idiid, Ann Chim Phas, 1894, [vn.], 2, 527, Seidell, Sdahilder (Coeby Lockwood & Son, 1907, p. 337. In table given, a=grains TINO<sub>2</sub> per 100 grains of solution, and b=grains TINO<sub>2</sub> per 100 grains of water.

2 Retgers, Jahib. Mr. 1, 1893, 1-90, 1896, ii. 183, Penfeld and Kreider, Amer. J. Sci. 1804 (m.) 48, 141

Reigers, Jahrb. Mit., 1893, 1-90, 1896, II. 183, Fenned and Kreiser, Amer. J. 1894, [III.], 48, 111
 III. L. Wells and Metzer, Amer. Chem. J., 1901, 26, 271, cf. Date, Compt. rend., 1879, 89, 575, 641
 Van Eyk, Zeitsch. physikal. Chem., 1905, 51, 721. Chem. News, 1905, 91, 295.
 Fock, Zeitsch. Krysl. Men., 1889, 4, 583, 1897, 28, 337
 Van Lyk, Proc. K. Akad. Webasch. Amsterdam, 1899, 1, 229. Zeitsch. physikal. Chem. 1899, 39, 430, 1905, 51, 721. Chem. News, 1905, 91, 229. Zeitsch. physikal. Bull. Soc. Juna. Min., 1905, 28, 311, Gossnet, Zeitsch. Kryst. Min., 1903, 38, 110, 3tortenbeker, Zeitsch. physikal. Chem., 1903, 43, 629
 Wallerant, loc. cit.

Wallerant, loc. cit., Compt. rend., 1905, 140, 1045.



Fro, 18 - Equilibrium diege au for the system thallons nitrate-polassium nitrate.

Silver nitrate forms no mixed crystals with thallous nitrate, but forms a compound, TINO, AgNO, which melts at 82.8°, and only separates as the initial solid phase from fused mixtures containing 48 to 52 molecules per cent of thallons nitrate.

One or two other double intrates are known.2

Thallic nitrate, Tl(NO<sub>3</sub>), is prepared by dissolving thallic oxide in nitric acid. It separates from an acid solution in large, colourless, transparent, deliquescent crystals which contain 311,0.3 The salt is readily hydrolysed by water, and is easily decomposed by heat

The double salt potassium thallie nitrate, K2Tl(NO3), H2O, readily crystallises from a solution of the mixed mitrates in intric acid (R J. Meyer)

Thallothallic nitrate, Tl(NO<sub>3</sub>), 2TINO<sub>3</sub>, is readily obtained by dissolving thallous intrate in warm intric acid of density 150. The double salt separates in prisms, melts at 150°, and is stable in dry air

Thallous hypophosphite, TIII.PO, is a sparingly soluble salt which crystallises in rhombic bipyramids (a/b = 0.786/1/0.806). It melts at 150, and at higher temperatures decomposes, evolving phosphine and leaving a residue of thallous meta- and pyro phosphates 5

Thallous hypophosphate, Tl<sub>1</sub>P<sub>2</sub>O<sub>6</sub>, from thallous carbonate and hypo phosphoric acid, is a sparingly soluble salt, which turns blue when exposed to light, and decomposes at 250° into thallnun and thallnun metaphosphate.

The acid hypophosphate, Tl,H,P,O,, prepared from the acid barium salt and thallons sulphate, forms monochnic crystals, slightly

soluble in cold, more readily in hot water. It melts at 200° and evolves phosphine at higher temperatures

Trithallous orthophosphate, Tl PO4, is obtained by adding excess of thallous carbonate to phosphorie acid, by mixing thallous sulphate and disodium hydrogen phosphate and adding a little ammoma, or by fusing Unum metaphosphate with two molecular proportions of sodium carbonate.

- Hises in long, colourless, anhydrous needles of density 6.89, and 1 Dennis, Dennis, J heat One litre of a saturated aqueous solution contains 4 97 Willin, Castangibate at 15° and 6.71 at 100°. The phosphate is insoluble
- Carstanjen, J. prakt .

  Carstanjen, loc cit., On.

  Vogel, Zeitsch anner Ched. Websich Amsterdam, 1900, 2, 480, Zeitsch, physikal.

Voxel, Zeitsch anner Ched. II desich Amsterdam, 1900, 2, 480, Zeitsch, physikal.
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 Przibylla, Zeitsch anner, Ca and Wendel, E. r. 1903, 36, 4055.
 1913, 103, 2116
 35, 207; R. J. Meyer, Zeitsch anner Chem., 1900, 24, Europe Land, Ann. Chem. Phys., 18t 311,0
 [iv], 17, 310, Miller, Proc. Koy Schem. J., 1901, 26, 275.
 Van Eyk, Zeitsch physikal. (Ni2
 Kryst Min., 1903, 38, 110, Reigers, 649; Rummelsberg, J. makt. Chem., 1892, [11.], 10 Thomas, Compl. rend., 1904, 138,

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in alcohol, but dissolves readily in aqueous solutions of ammonium salts. The aqueous solution has an alkaline reaction 1

Dithallous hydrogen phosphate, TI, HPO, H2O, may be prepared by neutralising a boiling solution of phosphone and with thallous carbonate, evaporating the liquid to a syrup, and allowing it to crystallise. The crystals are orthorhombic (a b c=0.934 1 0.657) and isomorphous with sodium dihydrogen phosphate, NaH<sub>2</sub>PO<sub>4</sub> H<sub>2</sub>O.<sup>2</sup> The anhydrous salt has a density of

Thallous dihydrogen phosphate, TlH2PO4, is obtained by adding phosphoric acid to a solution of the foregoing salt and crystallising the solution It crystallises in pearly plates or long needles, which are monoclinic (a b c=3.175 1 1.458;  $\beta=91^{\circ}$  14') and isomorphous with disammonium hydrogen phosphate, (NH<sub>4</sub>),HPO<sub>4</sub> The salt melts at 190°; its density is 4 723 The aqueous solution has an acid reaction 4 According to Lamy and Descloreany, there exists a double salt of the formula TLHPO, 2THLPO,

Thallous pyrophosphate, Tl<sub>1</sub>P<sub>2</sub>O<sub>5</sub> is prepared by heating dithalium hydrogen phosphate to dull redness. It dissolves in water (with the separation of some insoluble white powder), and on evaporating the solution, beautiful, lustrous monoclinic crystals (a b c - 1:127 + 1:2921 +  $\beta$  - 111' 0') of the anhydrous pyrophosphate separate. On standing, the mother liquor deposits monoclinic crystals ( $a \cdot b \cdot c - 2.1022 \cdot 1 \cdot 1.9217$ ,  $\beta = 11.1^{\circ} 57'$ ) of the dilividrate, Tl<sub>1</sub>P<sub>2</sub>O<sub>7</sub> 2H<sub>2</sub>O The density of the anhydrous salt is 6.786

In acid pyrophosphate, TLH<sub>2</sub>P<sub>2</sub>O<sub>3</sub>, is obtained by heating dihydrogen thallum phosphate to 210° It is readily soluble in water, from which it crystallises in small, short prisms, melts at 270, and is converted into the metaphosphate at a red heat "

Thallous metaphosphate. TPO, is obtained as a white solid, sparingly soluble in water, by heating dilivdrogen thallium phosphate to reduces. A more soluble modification is said to be formed by heating drammonnim thallium phosphate?

Thallous thiophosphate, TLPS, prepared by heating thallons sulphide with prosphorus pentasulphiele, is a vellow, crystalline solid, insoluble in water, alcohol, ether, benzene, and carbon disulphide.

Thallic orthophosphate, TIPO, 211.0, is said by Willin to be obtained by adding phosphoric acid to concentrated thallic nitrate solution, and then adding water. It is described as a white, very sparingly soluble substance, readily hydrolysed by boiling water 9

Thallous orthoarsenite, TlAsO, is a yellow, crystalline solid, sparingly soluble in water, obtained by boiling a solution of thallous sulphate with potassium arsenite and a little potassium hydroxide  $^{19}\,$ 

Crookes, Willin, Lamy, Ann. Chem. Phys., 1805, [w], 5, 119. Ball. Soc. chem.,
 1869, [u.], 11, 210. Rammelsberg, Ber., 1870, 3, 276, 1882, 15, 2228. Poor Annalen,
 1872, 146, 592
 Rammelsberg, Ioc. cd., Lamy, Ioc. cd., Lamy and Desclore ux, Ann. Chem. Phys.,

Ramm (sterg, toc. cit., namy, tow. cit., namy, 1869. [1v.], 17, 310

Schroder, Ber., 1871, 7, 676

Rammelsberg, toc. cit., Lamy and Desclotzeaux, toc. cit., Lamy, toc. cit., Lamy, and the clotzeaux, toc. cit., Lamy, toc. cit., Brand, Zeitsch. and, Chem., 1869, 28, 595

Lamy, loc cit
 Clamy, loc cit
 Clattel, Zetsch anong Chem, 1893, 4, 186
 Willin, Ann. Chem. Phys., 1865, [vv], 5, 5.
 Stavenhagen, J. prakt. Chem., 1895, [ii], 51, 1.

Trithallous orthoarsenate, Tl, AsO, is obtained as a white, crystalline precipitate by adding ammonia to an aqueous solution of thallium dihydrogen arsenate 1

Dithallous hydrogen arsenate, TlallAsO, is obtained by neutralising a boiling solution of atsenic acid with thallous carbonate and cooling the solution I It forms long, transparent needles, readily soluble in water. When strongly heated it gives oil water, and then decomposes, leaving a black residue of thallie oxide (Lamy)

Thallous dihydrogen arsenate, TIII, 4sO,, was prepared by Willin by boiling a solution of aisemons acid with thallic oxide, concentrating, and crystallising. It forms gittering, colourless needles and is readily soluble in water.1

Thallic orthoarsenate, TIASO, 2112O, has been described by Willin 1 Thallous metantimonate, TISbO, 211,0 has been described by Berlstein and Blase 2

Thallous vanadates (see Vol. VI. of this series)

### THALITIM AND THE CARBON GROUP,

Thallous carbonate, Tl2CO is obtained by passing curbon dioxide into thallous hydroxide solution, by exposing granulated thalloun to warm air for some time and their heating it with ammonium carbonate solution, or by shaking and heating barroni carbonate with thallous sulphate solution. The carbonate crystallises from water in colourless, glittering, monochine prisms (a b c -1 3956 | 1 19586 ,  $\beta$  =94° 47') of specific gravity 7 164,3 melts at 273°,1 and at high temperatures loses carbon dioxide. The carbonate is insoluble in alcohol, 100 grains of water dissolve the following amounts of the salt (Lamy, Crookes) -

The solution has a caustic taste and an alkaline reaction.

An acid carbonate is not known in the pure state, in spite of various attempts to prepare it 5. A basic carbonate has been described

Thallous cyanide, TICN, may be obtained by adding thallous hydroxide to prussic acid, or by mixing concentrated solutions of thallous intrate and potassium cyanide, when thallous examile is precipitated? It is best prepared from barum cyanide and thallous sulphate. It crystallises from water, in which it is not very soluble, in glittering plates. When heated it decrepitates, melts, and decomposes

One hundred grams of water dissolve 8 67 at 0°, 15 17 at 14°, and 29 57 grams at 31°, of thallous cyanide. The solution reacts alkaline and smells of prussie acid. The cyanide is soluble in alcohol, but very sparingly soluble

Belistein and Blase, Chem. Zentr., 1889, 803
 Lamy and Descloreaux, Ann. Chem. Phys., 1869, [iv.], 17, 310, Millet, Proc.

Roy. Soc., 1865, 14 556

Carnelley, Trans. Chem. Soc., 1878, 33, 273.

Cartainen, J. publ. Chem., 1867, 102, 65, 129., Giorgia, Gazzetta, 1891, 24, ii. 474; Att. R. Acad. Lince, 1894, [v.], 3, ii. 104

Wyroubolf, Bull. Soc. trang. Mer., 1889, 12, 136.

<sup>\*</sup> Crookes . Kuhlmann, Compt rend., 1862, 55, 607.

THALLIUM. 19

in a mixture of alcohol and ether. It dissolves in aqueous potassiun eyanide, forming soluble potassium thallium cyanide, and forms minieron double and complex eyanides with other cyanides.1

Thallothallic cyanide. TICN TI(CN), is prepared by dissolving thallicoxide in aqueous prinsic acid, and crystallises from the solution in colour less, orthorhombic crystals One hundred grams of water dissolve 9.75, 15-29 and 27.31 of the salt at 0', 12°, and 30° respectively, giving a neutral solu tion in which the salt is probably dissociated into the ions Tl and Tl(CN)4'.1 From the cold, aqueous solution, alkalies precipitate thallie hydroxide, but no precipitate is produced by aminonia. At 125° to 130° the salt melts and decomposes 3

Thallous cyanate, TICNO, precipitated in small, glittering crystals by mixing alcoholic solutions of thallons acetate and potassium cyanate, is very soluble in water 1

Thallous thiocyanate, TICNS, is precipitated as small, white, tetranamous thiocyanate, (1935), is precipitated as sman, white, section good crystals (a/c) 1/0/5593) when agreed solutions of thallons intrate and potassium thiocyanate are mixed? One hundred grains of water dissolve 0/316, 0/392, and 0/732 grains of the salt at 19/94\*, 25°, and 39/75° respectively 6

The double salt, TICNS KCNS, crystallises from a solution of thallous and potassium thiocyanates 7. The compound, Tl<sub>2</sub>Pt(CNS), 9NH<sub>39</sub> is also

Thallous oxalate, Tl.C.O. prepared from thallous carbonate and oxalic acid, crystallises from boiling water, in which it is not very soluble, in small, pearly, monoclime prisms (a |b|c=1 1384 | 1 | 2 2405 ,  $\beta=99^\circ$  137) of density 6.31. One little of a saturated aqueous solution of the salt contains 15.77 grams of the oxalate at 20°, and 18.69 at 25°. Thallium hydrogen oxalate, THC,O, also crystallises in monochiuc prisms

Thallic oxalate,  $\text{TL}(C,O_1)_p$  is precipitated when alcoholic solutions of thallic formate and oxalic acid are mixed. It is not very stable. Thallic hydrogen oxalate,  $TIH(C,O_4)$ , 3H,  $O_7$  is readily obtained as a white, sparingly soluble, crystalline powder, by the action of aqueous ovalre act at 25° on thallic hydroxide or an acetic acid solution of thallic acetate. The corresponding potassium and ammomum salts may be obtained by precipitating acid solutions of thallie salts with potassium and ammonium oxalate respectively More complex ovalates are also known, and also thallothallic oxalate, Tl (C2O1), 311,0 10

<sup>&</sup>lt;sup>1</sup> Fronmuller, Ber., 1873, 6, 1178, 1878, 11, 91, Fischer and Benzian, Chem. Zeit. 1902, 26, 19. Cunningham and F. M. Perkin, Trans. Chem. Soc., 1909, 95, 1509. For terro-, ferri, and coball equandes, see Vol. IX.
<sup>2</sup> Goldschmidt, Inaugural Dissertation (Berlin, 1903).
<sup>3</sup> Fronmuller Leave.

<sup>3</sup> Fronmuller, loc cut

<sup>&</sup>lt;sup>2</sup> Fronmuller, for cet

<sup>4</sup> Kuhlmann, Compt rend, 1802, 55, 607

<sup>5</sup> Miller, Proc Roy Soc, 1805, 14, 555, Phil Mag, 1866, [iv], 31, 153,

<sup>6</sup> Bottger, Zeitsch physical Chem, 1903, 46, 602, A.A. Noyes, vind, 1890, 6, 249;

Noyes and Abbat, sbid., 1895, 16, 132.

<sup>7</sup> Carstangen, J. prakt Chem, 1867, 102, 65, 129.

<sup>8</sup> Peters, Zeitsch anong Chem, 1912, 77, 137

<sup>8</sup> Lanny and Peccloreaux, Ann. Chem Phys., 1869, [iv], 17, 310, Abegg and Spencer,
Zeitsch anong Chem, 1905, 46, 406

<sup>9</sup> Rabe and Stemmetz, Ber., 1902, 35, 4447, Zeitsch anong Chem, 1903, 37, 88;

R. J. Meven and Goldschmidt, Ber., 1903, 36, 138

Abegg and Spencer, Zeitsch, anong, Chem, 1905, 44, 379, 46, 406, Alugg and Schaefer, Jud., 1905, 44, 379, 46, 406, Alugg and Schaefer, Jud., 1905, 45, 283, Strecker, Annalen, 1865, 135, 207.

Thallium alkyl compounds.—Thallium compounds analogous to the aluminium tri-alkyls have not yet been prepared, but numerous compounds of the type Th(Alkyl),X are known. They must be looked upon as binary electrolytes, which dissociate thus

$$Tl(Alkyl)_2 \setminus --- Tl(Alkyl)_2 + X'$$

a view in accordance with the conductivities of their solutions, and with the fact that they are quite stable towards boiling aqueous aminonia and alkali hydroxides

The thallium alkyl chlorides are readily prepared by the general reaction .--

$$TICL_1 + 2Mg Mkyl Cl = TI(Alkyl)_0Cl + 2MgCl_0$$

the reagents being mixed in dry ethereal solution. The broundes, iodides, carbonates, etc., may then be prepared by double decomposition. The salts crystallise well, and are only sparingly soluble in water The hydroxides are strong bases, and reachly absorb carbon dioxide. The alkyl compounds thus bear a great resemblance to the thallous compounds 1

Thallous Alcoholates. -- Anhydrous thallous oxide dissolves in absolute alcohol, producing an alcoholic solution of thallous ethoxide, C<sub>2</sub>H<sub>5</sub>O Tl. A better method of preparation consists in suspending very thin thallium foil over absolute alcohol in an apparatus through which pure, dry oxygen is slowly passed. Even at ordinary temperatures the thallium rapidly disappears. The alcoholic solution thus obtained is heated to 100° to remove excess of alcohol, and thallous ethoxide is thereby obtained as a yellow, only liquid. The density of the liquid is 3 522 at 20°, and 3 562 at 0°, so that it is an extremely heavy liquid. The refractive index for sodium light, 1 6826 at 20°, is also very high. Thallous ethoride is readily hydrolysed by water to thallous hydroxide and ethyl alcohol. The corresponding amyl derivative is an only liquid, but the methyl compound is a solid the specific gravity of which is about five?

Thallous acetylacetonate, (CII, CO), CH Tl, prepared by heating thallous hydroxide with an alcoholic solution of acetylacetone, crystallises in flat, colourless needles, and melts with decomposition at 153°-160° 3

Other organic derivatives containing thallium cannot be described here, but the appended references to the literature may be of use 1

Thallous Silicates .-- Thallous oxide or carbonate reacts with silicite acid after the manner of the corresponding sodium and potassium compounds 5 A crystalline silicate of the composition 3TLO 2SiO, ILO has been described by Wyrouboff.6

<sup>&</sup>lt;sup>1</sup> R. J. Meyer and Bettheum, Ber., 1904, 37, 2051; Shukoll, Ber., 1905, 38, 2691; Harsen, Ber., 1870, 3, 9, Hartwig, Ber., 1874, 7, 298, Annaten, 1875, 176–257. Canna and Frommuller, Ber., 1874, 7, 302.

<sup>a</sup> Lamy, Ina Chim. Phys., 1864, [Iv.], 3, 373; Kahlbaum, Roth, and Siedler, Zeitsch. anoig Chem., 1902, 29, 177.

<sup>a</sup> Kurowski, Ber., 1910, 43, 1078, Morgan and Moss, Trans Chem. Soc. 1914, 105, 189.

<sup>a</sup> Kuhlmann, Compt. rend., 1862, 56, 607, Ana. Chim. Phys., 1863, [in.], 67, 431, Bull. Soc. chim., 1864, [ii.], I. 330, de la Provostave, Compt. rend., 1862, 56, 610, Crookes, Chem. News, 1864, [ii.], I. 330, de la Provostave, Compt. rend., 1862, 56, 610, Crookes, Chem. News, 1864, [ii.], J. J. Chem. Soc., 1864, 17, 112. Willin. Ann. Chim. Phys., 1865, [iv.], 5, 5, Lamy and Desclotzeaux, ibid., 1869, [iv.], 17, 110. Lescuru, Bull. Soc. chim., 1875, 24, 516. R. J. Meyer, Zeitsch. anoig. Chem. 1900, 24, 321, Rabe and Steumetz, ibid., 1903, 37, 88. Hawley, J. Amer. Chem. Soc., 1907, 29, 300.

<sup>b</sup> Flemming Johnesber., 1868, p. 250.

<sup>e</sup> Wyronboll, Bull. Soc. franc. Min., 1889, 12, 536.

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#### THALLIUM.

Thallium enters into the composition of various kinds of optical glass of high refractive index. High retractive power is a general characteristic of thalloun salts (pp. 189, 190, 192, 200).

Thallous silicofluoride, Tl SiF6 is prepared from thallons eubonate and hydrofluosilicie acid, and crystallises from aqueous solution in colourless octahedra According to Kuhlmann, the salt is volatile without decomposi tion, and crystallises from aqueous solution in hydrated form 1

## THAILIUM AND BORON

Thallous Borates —Thallons oxide dissolves in fused boron sesqui oxide, and the liquid mass solidities either to a clear glass or a crystalline solid 2 Various borates of thallum have been described by Buchtala !

Thallous tetraborate, Tl<sub>2</sub>B<sub>1</sub>O<sub>7</sub> 2H<sub>2</sub>O<sub>7</sub> is obtained by dissolving thallous carbonate (1 mol.) and bone acid (1 4 mols) in water and crystallising. If is also obtained, together with thallous hexaborate, Tl,B,O<sub>10</sub> 311,O, by fusing the carbonate (1 mol.) with bone acid (6 mols.) and existallising the product from water. If more bone acid (8 mols.) is used, a mixture of the hevalor ite and thallous decaborate,  ${\rm Tl_2B_{10}O_{11}~8H~O_{18}}$  is produced

Thallous octaborate. Tl B<sub>1</sub>O<sub>11</sub>4H O<sub>1</sub> and thallous dodecaborate,  $TL_iB_{12}O_{16}(7H_2O_i)$  are prepared by dissolving thallous carbonate (1 mol.) in excess of aqueous born and (10 mols). The dode aborate also forms a pentalized attention that is monoclime (a  $b \in [1.583]$  1 1 955  $\beta = 91^{\circ}$  25') 4

Thallous metaborate, 211BO/11/O, crystallises from water in coloniless, wedge shaped crystals which turn roll and black on exposure to an

Thallous perborate, 
$$^3$$
 Tl<sub>2</sub>B<sub>2</sub>O<sub>7</sub>,  $re$   $^{-1}$  TlO B O O, is obtained as a white TlO B O O

powder by adding 30 per cent. hydrogen peroxide to an aqueous solution of any of the preceding borates—It shows the usual reactions of a per salt and explodes when rapidly heated—The aqueous solution decomposes on evaporation, thallous metaborate being produced

# DETECTION AND ESTIMATION OF TRALLICM

The compounds of thalling impart a green colour to the Bursen flame, and pectroscopic examination of the flame reveals the green line  $\lambda$  5350-7. Thallium compounds are poisonous

The following are the clief reactions of thallows salts -(1) Hydrogen sulphide incompletely precipitates thallium as the brown sulphide, insoluble in acetic acid. In the presence of mineral acid no precipitate is obtained (n) Ammonium sulphide leads to complete precipitation of the thallium as sulphide. (iii) Hydrochlone acid or a soluble chlonde precipitates the thallium almost completely as white thallous chloride (iv) A soluble iodide precipitates thallimin completely as yellow thallous iodide (v) Potassium

Kuhlmann, Bull. Soc. chim., 1864, [n.], 1, 330., Compt. cend., 1864, 58, 1037.
 Weither, J. piakt. Chem., 1864, 92, 128.
 Guertlei, Zeituh. anning Chem., 1964, 40, 225.
 Buchtala. I. piakt. Chim., 1913 [n.], 88, 771.
 Scharizei, Zeitsch. Kryst. Min., 1914 54, 232.

chromate in neutral or ammoniacal solution precipitates thallium quantitatively as the yellow chromate (vi) Chloroplatinic and precipitates thallium quantitatively as yellow thallous chloroplatinate (vii) Sodium cobaltimitrite precipitates thalloun quantitatively as scarlet thallous cobaltinitrite (viii) Potassium iodide and antimony trichloride lead to the precipitation of orange red thallous antimonious iodide, 3TH,2SbL,2

Thallie salts give a brown prespitate of Tl(OH), with alkali (including ammonium) hydroxides, insoluble in excess of reagent. With a soluble iodide they give thallous iodide and iodine. Alkali chloride or chromate produces no precipitate

The detection of thallium in qualitative analysis has been studied by A A Noyes, Bray, and Spear, to whose work the reader is referred for details 3

The gravimetric estimation of thallium is somewhat difficult, since its "msoluble" salts usually have a perceptible solubility, are difficult to wash, and are distinctly volatile at high temperatures. The usual method is to precipitate as thallous iodide at 80° to 90°, wash with 80 per cent alcohol, dry at 170, and weigh! Thallium may also be estimated as thallous chloroplatmate, I part of which dissolves in 15,600 of water at 15°, and in 1950 at 100° 5. It may be precipitated and weighed as thallors chromate. or by evaporation with sulphune and it may be converted into either thallous hydrogen sulphate at 220° to 230° or thallous sulphate at a dull red heat? Thillium may also be precipitated as thillons thiostamate,  ${\rm Tl_1SnS_n}$  dried at 105°, and weighed. Thillium salts may be oxidised to thallic salts with brom unic acid, and estimated by weighing the precipitated gold 9

The electrolytic determination of thallimm is not a very accurate or convenient process, and is best effected by depositing the thallmin into a mercury cathode, or, better, a dilute zinc or cadminim amalgam 19

Several volumetric methods for the estimation of thallium have been proposed. Thallous salts in the presence of hydrochloric acid may be oxidised to thallie salts by standard permanganate, 11 or they may be heated with potassium brounde, dilute hydrochloric acid, and an excess of standard

Cunningham and Porkin, Teans Chem Soc., 1909, 95, 1569. Tanatar and Petroff, J. Russ. Phys. Chem. Soc., 1910, 42, 94
 Ephraim Zertsch annug Chem., 1908, 58, 353
 A. A. Noves, Bray, and Speat. J. Amer. Chem. Soc., 1908, 30, 481. Chem. News, 1908, 98, 6, etc., see also Sorby, Chem. Lens, 1869, 19, 309, Ross, Chem. Zinte, 1832, p. 54, Behrens, Zertsch. anal. Chem., 1891, 30, 138. Chapman, Phil. Mag., 1876, [v.], 2, 397. Phipson Cound rend., 1874, 78, 563
 Baubiguv, Compl. rend., 1874, 78, 563
 Baubiguv, Compl. rend., 1891, 113, 514. Weither, Zertsch. anal. Chem., 1861, 3, 1;
 Long, thid., 1891, 30, 342. Hebberling, Annalen, 1865, 134, 11
 Crosbes, Chem. Mices, 1864, 9, 1. Neumann, Annalen, 1888. 244, 319, Cushman, Amer. Chem. J., 1900, 24, 222
 Browning and Hutchius, Amer. J. Sci., 1899, [iv.], 8, 460. Chem. News, 1899, 30, 286

<sup>\*</sup> Howning and Filterian, Amer. Chem. Soc., 1900, [11.], 9, 137, Zeitsch anorg Chem., 1900, 23, 155

\* Hawley J Amer Chem. Soc., 1907, 29, 1011

\* Thomas, Compit. rend., 1900, 130, 1316, Bull Soc. chem., 1902, [11.], 27, 470;

Ann Chim Phys., 1907, [viii.], 11, 204.

10 Morden, J Amer. Chem. Soc., 1909, 31, 1045; of Neumann, Ber., 1888, 21, 356;

Schueht, Zeitsch. anal Chem., 1883, 22, 241. Heiberg, Zeitsch anorg. Chem., 1903, or 347.

 <sup>35, 847.
 35, 847.</sup> Willin, Ann. Chim. Phys., 1865, [iv], 5, 5. Hawley J. Amer. Chem. Sur., 1907,
 29, 800, Λ. A. Noyes, Zeitsch phesikal Chem., 1892, 9, 603.

potassium bromate, the bromme distilled into potassium iodide and the liberated iodine titrated with thiosulphate. The iodine found is a measure of the oxidising power of the bromate present in excess of that required for the reaction —

$$3TICI + NaBrO_3 + 6HCI + 3TICI_1 + NaBr + 3H_2O$$

Lead is reachly separated from thallium either by precipitation as the sulphide in dilute immeral acid solution, or by evaporation with sulphiric acid to remove the lead as the insoluble sulphate. Silver may be separated from thallum by passing a rapid stream of chlorine through the solution. The silver is precipitated as chloride, the thallum remaining in solution as thallie chloride.

Marshall J Soc Chem Ind., 1909, 19 991. For other methods, vide Fert Zeitsch anal Chem., 1889, 28, 314.
 Spanholz duci. 1893, 31, 519.
 Cannet, Compt. rend., 1889, 109, 177.
 Nuctski, Dingl. pulm. J. 1876, 219, 262.
 Thomas, Compt. rend., 1902, 134, 655.
 Inn. Chem. Phys., 1907 [vm.], 11, 291.
 Bowning and Pilmer, J. Sec., J. 1909, 191.
 Sie also Weiner, Chem. New, 1886, 53, 51.
 Spencer and Le Ph. Trans. Chem. Soc., 1908, 93, 858.
 For intrinsic uniformation on the analytical chemistry of the Physics.
 Crookes, Select Methods of Chemical Analysis, (Longmans & Co., 4th ed., 1905).

## CHAPTER IX.

# SCANDIUM.

Symbol, Se Atomic weight, 11.0 (O = 16).

Occurrence - Scandium, despite its rarity, is extremely widely diffused in nature. In almost all rocks from which the mountains of the earth, or rather the chief parts of the earth's crust are formed, semidium can be detected, as the comprehensive spectroscopic researches of Eberhard <sup>1</sup> have shown.

The more commonly occurring immerals which contain scandium are euxenite2 and keilhauite (p. 221) from certain localities, mica from Ytterby, and cossiterite and wolframite from the mountain ranges of Saxony and Bohemm, especially from Zimiwald in Saxony 1. They are all derived from gramtes or certain pegmatites 1

It will be noticed that scandium occurs in certain rare earth immerals. However, it is by no means an invariable companion of the rare earth elements. In the course of some ten years of work on the rare earths, Urbain never met with scandium in any of his fractions,3 and Hicks failed to detect scandium in specimens of fergusonite, aschinite, cusenite, and samarskite examined by him 6. Scandhum occurs in gadolinite, but only in the deposits found at Ytterby. When scandium occurs in association with the rare earth elements, the yttrium group usually predominates, and hence the occurrence of scandium in Finnish orthite, a cernon nineral, is of interest 7

The occurrence of scandium and the rare earth elements in the rare numeral with ite was discovered by Crookes,8 who isolated considerable quantities of scandia from this source. Withit is a black, absorphous mineral found with monazite in a felspar quarry at Impilaks, Lake Ludoga, Finland. Density, 4.85., hardness, 6. The mineral is feebly radioactive,

Eberhard, Sitzungsber, K. Akad. Wiss. Berlin, 1908, No. 38, p. 851, 1910, No. 22,
 404; Chem. News, 1909, 99, 30, 1910, 102, 211.
 See p. 220; Hauser and Wirth, Ber., 1909, 42, 4445.
 Seandown has been found in American wolframite. Lukens, J. Amer. Chem. Soc.,

Scandium has been found in American wolfiamite. Lukens, J. Amer. Chem. Soc., 1913, 35, 1470.
 Eberhard, Loc ett., Vernadski, Bull. Acad. Sci. Petrograd, 1908, p. 1273
 Urbani, J. Chem. phys., 1905, 4, 64, cf. Crookes, Chem. News, 1905, 91, 61, Eberhard, Zeitsch, anory Chem. 1905, 45, 374.
 Hicks, J. Amer. Chem. Soc., 1911, 33, 1492
 R. J. Mayer, Sitzungsber, K. Akad. Wiss. Berlin, 1911, p. 379.
 Crookes, Phys. Tenn. 1, 1908, A, 209, 15. Proc. Roy. Soc., 1908, A, 80, 516; Chem. News. 1908, 68, 224.

News, 1908, 98, 274.

and when heated to reduces gives off water, hydrogen sulphide, helium, a trace of neon, etc. The wikite examined by Crookes had approximately the following composition -

| Ta <sub>2</sub> O <sub>5</sub> and Cb <sub>2</sub> O <sub>5</sub><br>TrO <sub>2</sub> and ZrO <sub>3</sub> |   | 25 cent<br>15 91<br>23 36 | -   | FeO .<br>UO                |   | Per cent<br>15/52<br>3/56 |
|------------------------------------------------------------------------------------------------------------|---|---------------------------|-----|----------------------------|---|---------------------------|
| Ce <sub>2</sub> O <sub>3</sub>                                                                             |   | 2.55                      | -   | SiO,                       |   | 16 98                     |
| $Y_2O_3$                                                                                                   |   | 7.61                      | - 1 | H <sub>2</sub> O and gases | - | 5.83                      |
| Se <sub>2</sub> O <sub>3</sub><br>ThO                                                                      | • | 1 17<br>5 51              |     | Undetermined               |   | 1 97                      |

The particular deposit of this miner dused by Crookes is now exhausted, and the other deposits do not appear to contain more than the merest traces of scandium 1

The only known mineral in which scandium occurs as an essential constituent is the rare numeral therefore, (8cA) ScO This is a grevishgreen, translucent silicate of scandium and yttrimi which less been found in pegmatite at Iveland, South Norway - Hardness, 6 5, density, 3 57 - The inmeral is orthorhombic (a |b||c=0.7456 + 1/1912):

Scandium has been shown spectroscopically to occur in relative abundance in the sun and stars

History In 1879 Nilson was preparing atterbia from cuxemto and gadolimite by the procedure that had been given by Marignae, the discoverer of the earth (p. 225) . In so doing, he made the discovery of a new earth, present in very small quantity, and characterised by its very feeble basicity (less than that of ytterbia), its very low chemical equivalent, and its spark spectrum, Only 0.3 grain or the earth was obtained, and that in an impure state. To the new element present in the earth, Nilson gave the name scandium few weeks after the announcement of the discovery, Cleve reported that he had isolated 0.8 gram or scandia from 4 kilos of quidolimite, and 1.2 grams of scandia from 3 kilos of keilhanite. Cleve also described a few compounds of scandium (  $\Lambda$  little later, Nilson obtained a few grains of scandia, described several scandium salts, and determined the atomic weight o' the element  $\delta$ 

Scandnum corresponds very closely with ekaboron, one of the elements predicted by Mendelceff when he put forward the periodic classification of the elements, and its discovery, coming only four years after the discovery of eka-aluminium or gallium,4 was a matter of great interest and assisted largely in the recognition of the merits of the periodic classification by chemists in general. The identity of ekaboron and scandium was pointed out by Cleve, \$\infty\$ the following table will serve to show how closely the properties of scandium were predicted by Mendelceft - a

<sup>&</sup>lt;sup>1</sup> R. J. Meyer and Winder, Zerish anna Cuem., 1910, 67, 308. Chem. News, 1910. 102, 163

Scheelig, Mineral Zente, 1911, p. 721
 Scheelig, Mineral Zente, 1911, p. 721
 Silsson, Compt. rend., 1879, 88-645, 1880, 91, 56, 118. Ber., 1879, 12, 550, 554;
 1880, 13-14-10, 1139. Otters of K. Svenska Vet. Ikad. Fenhandt., 1879, No. 3, 1880,
 No. 6, Cleve, Bull. Som. ekom., 1879, [n.], 31, 486. Compt. cond., 1879-89, 119. Chem.
 News 1879-40, 159. Otters of K. Svenska Vet. Akad. Ferhandt., 1879, No. 7
 Scep. 113
 Cleve, Compt. rend., 1879, 89, 419. Bull. Sov. ekom., 1879, [n.], 31, 486. Chem.
 News. 1879, 40-159.

News, 1879, 40 159.

Mendelétil, J. Russ Chem. Soc., 1869. 1, 60, 1-71, 3, 47, Annal. a. Suppl., 1872, 8, 133, trans. in Chem. News, 1879, 40, 213, etc., 1880, 41, 2, etc.

| Ekaboron                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | Scandium                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |  |  |  |  |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--|--|--|--|
| Atoma weight, 14 Should give one exide, Eb <sub>2</sub> O <sub>4</sub> , density, 3.5, more basic than Al <sub>2</sub> O <sub>4</sub> , less basic than MgO, insoluble in alkalies. Carbonate should be insoluble in water, and probably be precipitated as a basic salt. The double sulphates with alk it sulphates will probably not be alonis. Anhydrous chlorid, EbCl <sub>3</sub> , should be more difficultly volatile than AlCl <sub>4</sub> . In aqueous solution it should hydrolyse easier than magnesium chloride. Ekalonon will probably not be discovered appetroscopically. | Atomic weight, 41. Seandium oxide, Sc.O <sub>2</sub> , density, 3-86, is more base than Al <sub>2</sub> O <sub>2</sub> and probably less than MgO, insoluble in alkalies. Seandium carbonate, insoluble in water, readily loss earbon dioxide Double sulphates are known, but they are not alums. Seandium chloride, SeCl <sub>3</sub> , begins to sublime appreciably at 850°. In aqueous solution it is diedlighly hydrolysed Seandium was not detected by means of its spectium |  |  |  |  |

Extraction of Scandia from Minerals.—The method employed by Grookes in extracting seandia from winkite consisted in first isolating the seandia and rare earths, and then separating the seandia from this mixture by the fractional decomposition of the intraces. To obtain the mixture of seandia and rare earths, the immetal (1 pt.) was finely powdered and fused in a clay crucible with potassium hydrogen sulphate (5 pts.). The melt was cooled, pawdered, and mixed with five times its weight of water, stirred for twelve hours, and then filtered through a linen liter. The filtrate was boiled with an excess of minionia when all the earths were precipitated together with zirconia, titaine acid, ferric hydroxide, etc.—The well-washed precipitate was heated with a slight excess of oxalic acid, and when cold the oxalates were filtered and washed. The oxalates were next converted into anhydrous sulphates, dissolved in cold water, filtered, and the aminoma and oxalic acid processes repeated. The oxalates were then well washed and calcined to oxides.

The extraction of scandia from wolframite has been very thoroughly studied by R. J. Meyer and others. Only the wolframite from Zimiwald (Z) and Sadisof (S) is said to be worth working up for scandium, analyses of the carefully picked mineral yield the following results —

| $MO^{\gamma}$ | FcO.  | MnO.  | $\mathrm{PbO} + \mathrm{SnO_2}$ | $T_1O_2 + T_{3/2}O_6$ . | CaO  | Rare<br>Earths |
|---------------|-------|-------|---------------------------------|-------------------------|------|----------------|
| (Z.) 75·41    | 934   | 14 00 | 0 18                            | 0.50                    | 0.55 | 0 15           |
| (S.) 73·47    | 15.13 | 9.81  | 0.47                            | 0.63                    | 051  | 0 20           |

Rather more than half of the "rare earths" present consists of scandia

The finely powdered wolframite is fused with about 2.3 times its weight of sodium carbonate, a little potassium intrate being also added. The melt is boiled with water and a little alcohol, and the brown mixture of oxides left (which contains all the scandium) is washed, dissolved in hydrochloric acid, and the silica present removed in the usual manner.

The oxide residues from tungsten factories, if good Zimiwald wolframite has been worked up, are more convenient to use as raw material than

<sup>1</sup> For an account of which, see p. 353.

wolframite itself, and contain 0.30 per cent of rare earths. One kilogram of the oxide residues is dissolved in 3 littes of raw hydrochloric acid, the powder being introduced into the boiling acid. After drawing off the deposit of silica and boiling it with dilute hydrochloric acid, the filtrates are united

To separate the scandium, the preceding solution is boiled, and 40 grams of sodium silicofinoride are added slowly with stirring. The boiling is continued for half an hour, when the scandium separates completely as a white, pasty precipitate of scandium fluoride (produced by decomposition of the silicofluoride). The precipitate is filtered on a Nutschen litter and boiled with very dilute hydrochloric and. The fluoride is converted into sulphate in a platinum dish, the scandium precipitated as hydroxide with aminonia, and converted into oxalate by boiling with oxalic acid. The oxalate when washed and ignited leaves a white residue of nearly pure scandium oxide, which should weigh 3.0 to 3.3 grams.

The small amount of yttimin earths present in the scandia is easily removed. The scandia is dissolved in hydrochloric acid, excess of the acid removed, and the scandium precipitated at 100° as the basic throughpute. The washed precipitate is decomposed by hydrochloric acid, and the precipitation as throughpute repeated. The washed precipitate, now quite free from yttimin earths, is decomposed by hydrochloric acid, and the scandium in the filtered liquid precipitated as oxalate. The only impurity now remaining is thorum.

It is not difficult to reduce the amount of thoria in scandia to 0.5 per cent This may be effected in several ways, one of the most satisfactory being to pour the neutral solution of the chlorides of scandium and thorium into a large excess of 20 per cent, sodium carbonate solution, and boil the solution, when secuclium sodium errhonate is precipitated, the bulk of the thorum remaining in solution. The final 0.5 per cent of thoria, however, is difficult to eliminate. It cannot, for instance, be precipitated as the peroxide, and, moreover, it cannot be detected in the scandia by its arc spectrum. A number of methods for its elimination are known, the first being to drop slowly a neutral solution of the scandium salt (contaminated with thornin) into a concentrated (10 to 20 per cent ) neutral solution of ammonium tartrate, and boil the clear solution so obtained with ammonia. Scandium ammonium tartrate is thus precipitated free from thorium, and is washed with dibite ammonium faitrate solution. The second method can only be advantageously employed on a small scale. It consists in precipitating the thoroun from a intrate solution containing much free intric acid by the addition of an excess of potassium iodate sufficiently great to co-piccipitate a portion of the scandium. The liquid is then quite free from thorium. The third method consists in pouring a neutral chloride solution of the scandium cbloride into an execss of aqueous ammonium fluoride contained in a platinum dish and vigorously stirred. One gram of scandia requires 8 grams of animomium fluoride. The liquid is evaporated at 100°, and the thorium separates out as the insoluble fluoride, any scandium ammonium fluoride that crystallises

<sup>1</sup> R. J. Meyer, Zeitsch. anora. Chem., 1908, 60, 134, Chem. News, 1909, 99, 85, 97; D.R.P., 202, 523 and 208, 355. The method of extracting scanda from these oxide residues is also discussed in detail by Sterba Bohm (Zeitsch. Elektrochem., 1914, 20, 289)

See p 211.
 See Vol. V., and this vol., p. 320.

out may be redissolved by warming with more water, and the liquid then filtered. A fourth method, which is not, however, very convenient, is to separate the anhydrous chlorides of thorium and scandium by fractional sublimation, the former being the more volatile 1

Atomic Weight. - The analogy, though imperfect, between scandium compounds and the compounds of the rare earth elements, points to the tervalency of scandium, and hence, since the combining weight of scandium is 117, to the value 111 for the atomic weight. The specific heats of the oxide and sulphate may be cited in favour of this view, as the following comparison of molecular heats will show = 2

| $\Lambda l_2 O_3$ .              |  | 190  | $Al_{g}(SO_{1})_{3}$   |  | 63.6 |
|----------------------------------|--|------|------------------------|--|------|
| $G_{i} Q_{g}$                    |  | 19.5 | $Ga_2(SO_1)_3$         |  | 61.9 |
| ln <sub>z</sub> O <sub>τ</sub> . |  | 222  | In. (SO <sub>1</sub> ) |  | 66.4 |
| $S_{C_i}O_{i_0}$ .               |  | 20.8 | Sc.(SO.).              |  | 62.4 |
| Yt <sub>2</sub> O                |  | 23.3 | Yt.(SO <sub>1</sub> ), |  | 616  |

If the formula ScO be assumed for the oxide, then the specific heat of scandium works out to the abrormally low value of 3.95 between 0 and 100, if that of oxygen be taken as 30 (which is certainly not too great). The molecular weight of scandium acetylacetonate (p. 214) argues strongly for the tervalency of scandium, as also does the isomorphism of scandium ethylsulphate and acetylacetonate with the corresponding indium salts 3. Of the multiples of the combining weight (117, 294, 114, 588, etc.) which might be taken for the atomic weight, only the value 11.1 gives scandium a position in the periodic table, and it then occupies the place of Mendelcell's "ekuboron"

The value Sc 44'I is due to Nilson, the value 154 obtained by Cleve is undoubtedly too high. Preliminary experiments by R. J. Meyer and Goldenberg gave the values 41:11, 11-11, 14-29, and 13-90 for the atomic weight of scandium. An accurate determination of the atomic weight is, however, lacking at present. The determinations hitberto made have been effected by the "synthetic" sulphate method described later (p. 240).

# COMPOUNDS OF SCANDIUM

Scandium salts derived from colourless acids are themselves colourless and devoid of absorption spectra. They are diamagnetic, and have a sweet, astringent taste. In aqueous solution they are perceptibly hydrolysed. This will be seen from the following values for the equivilent conductivities of scandium chloride and other chlorides, the abnormal increase, in the conductivity of scandium chloride with dilution from v=32 to v=1024

\_\_\_\_\_

R. J. Meyer and Winter, Zedsch among Chem., 1910, 67, 398., Chem. News, 1910,
 162, 163, 175.
 R. J. Meyer and Goldenberg, thet., 1912, 106, 13.
 R. J. Meyer and Wassynehrow Zedsch among Chem., 1914, 86, 257.
 Sterba-Bobin, toe cit.
 Allson and Pettersson, Compt. int. J. 1880, 91, 232.
 J. Berr, 1880, 13, 1459.
 Janger, Proc. K. Akad. Welensch. Amsterdam, 1914, 16, 1095.
 Rev. 1880, 120.

<sup>33. 342.
4</sup> Clove, Compt. rend., 1879, 89, 419, Nilson, abid., 1880, 91, 56, 118, Meyer and Goldenberg, low est

As compared with the chlorides of the raise earth elements.

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litres being due to the high ionic mobility of the hydrogen ions of the acid set free by hydrolysis -1

| į                  |                                |                                  |                                  |                                  |                                    |                                  |                                   |
|--------------------|--------------------------------|----------------------------------|----------------------------------|----------------------------------|------------------------------------|----------------------------------|-----------------------------------|
|                    |                                | Tem                              | perature                         | , 25" ("                         |                                    |                                  |                                   |
| i                  |                                | _                                |                                  | -                                |                                    |                                  |                                   |
| r=                 | 1                              | 61                               | 128                              | 256                              | 512                                | 1024                             | λ <sub>1 121</sub> λ <sub>2</sub> |
| Set l <sub>1</sub> | 194.7<br>10.8<br>197.6<br>99.3 | 111 7<br>112 1<br>114 4<br>196 9 | 119 0<br>118 0<br>121 3<br>114 1 | 126 2<br>123 4<br>126 8<br>128 8 | 1 11 6<br>127 3<br>1 11 1<br>131 0 | 142 5<br>131 5<br>135 2<br>138 0 | 37 8<br>27 7<br>27 6<br>38 1      |

The basic oxide scandia, Sc.O., from which the scandium salts are derived, is decidedly stronger than alamma. It is, however, weaker than any of the rare earths of the type R O

Scandium is often included among the rate earth elements. In doing so, however, it becomes dafficult to decide whether search as to be regarded as an earth of the common or of the yttrium group. So far as besicity of oxide and solubilities of salts are concerned, scandium resembles the yttrium group on the other hand, the solubility of its double potassium sulphate places it with the cerning group. It differs from both groups in not forming a sulphate,  $S^{\pm}(SO_4)/SH(O_5)$  and a intrate,  $Sc(NO_3)/6H(O_5)$  . Further, scandium has a marked tendency to form what are probably complex as distinguished from double salts, and in this and in other respects it shows a remarkable rescribbance to thorning. The view of the fact that the cerum and vttrium groups or rare earth elements are distinguished by their platinocyamides (see [c 272), it is of interest to note that scandium platenor yanide, Se. Pt(CN)1], forms hydrates both with 18H O and with 21H O C

A few salts of scandium were described by Vilson and Cleve, many others have since been prepared and described by Crookes, and by R. J. Meyer and his co-workers

### SCANDIUM AND THE FITTORING GROUP

Scandium fluoride, ScF, is obtained as a white powder, difficult to filter, by he ting scandia with aqueous hydrofluoric acid, or by precipitating

<sup>&</sup>lt;sup>1</sup> Meyer and Wassinchnew, Zedoli and g. Chem., 1911, 86, 271, et. Meyer and Hauser, Lee chem. in: Indias decesitions Fider (Stinigar) 1912, p. 31. The artahydrate of the schuate, howere, is known. Compare, for example, the properties of the avalates, sulphates, double sodium sulphates carbonates and double carbonates and activiar tomates. Techniques, vide intra., Orloff, Chem. Zed., 1912, 36, 1407., Tschrivinski, Zedsch. Kryst. Min., 1913, 52, 11. Kryst Men , 1913, 52, 11 5 Nilson, Compt , and

Kyst Mm, 1913, 52, 41

Nilson, Compt. read., 1879, 88, 645—1850, 91, 56, 118., Ee., 1879, 12, 550, 554;
1880, 13, 1430, 1430, Cleve, Compt. read., 1879, 89, 419, Endt. Sec. clem. 1879, [n.], 34, 456., Chem. News, 1879, 40, 159., Chookes, Phil. Trans., 1908, A, 200, 15, 1910, A, 210, 359, From Row So., 1908, A, 80, 516, 1910, A, 84, 79, Chem. News, 1908, 98, 274, etc., 1910, 102, 73, etc., R. I. Meyer and Winder Zeitsch among Chem., 1910, 67, 398, Chem. News, 1910, 102, 163. R. I. Meyer with A. Wasspirthnew, N. Drapper, and E. Bodlandert, Zeitsch among Chem., 1914, 86, 257. The last paper contains a very full account of the chemistry of scale tem. References in the subsequent pages of the text to Crookes, Meyer and Winter, Meyer and Wasspirthnew, etc., refer to the papers enumerated in this footnote.

a solution of a scandium salt with hydrofluoric acid, an alkalı fluoride, or fluosibile acid. The fluoride is fusible with difficulty in the blow pipe flame. It is very slightly soluble in hydrochloric acid, in which it resembles the thornm salt and differs from the salts of the rare earth metals (Crookes; Meyer and Wassquehnow),

Scandium fluoride, like zirconium fluoride, is soluble in solutions of the alkali fluorides Scandium ammonium fluoride, ScF, 3NH,F, is readily soluble in water, from which it crystallises in octahedra. It is decomposed by hydrochloric or dilute sulphuric acid with the precipitation of scandium fluoride or scandium aumounum sulphate. In aqueous solution it may be regarded as yielding chiefly the ions  $3\mathrm{NH}_1'$  and  $\mathrm{SeF}_n'''$ , since it does not give any precipitate when boiled with ammonia. Sodium or potassium hydroxides, however, precipitate scandium hydroxide The corresponding potassium and sodium salts, ScF 3KF and ScF, 3NaF, are known, they are less soluble in water and less "complex" than the ammonium selt (Meyer and Wasspachnow)

Scandium chloride, SeCl, may be prepared by heating scandia in a current of disulphur dichloride vapour and chlorine. It is a white solid which begins to sublime at 800°. It is hygioscopic and dissolves in water with evolution of heat, but it is insoluble in alcohol (Meyer and Winter)

When scandium hydroxide or oxide is dissolved in hydrochloric acid and the solution concentrated, fine white needles of the hexahydrate, ScCl 6H,O, separate out. It is deliquescent and soluble in alcohol, dired at 100 it becomes converted into 2SeCl, 311 O, and this loses hydrogen collored when further heated (Crookes) An aqueous solution of scandium chloride reacts acid, owing to hydrolysis (p. 209)

Seanchum chloride forms a very soluble double chloride with custom chloride (Meyer and Wassjuchnow) The aurichloride, "SeCl. 2AuCl. 2111,0, forms readily soluble, yellow crystals, which lose water when carefully dried, and yield the hydrates with 8H2O and 2H2O, and finally the anhydrous salt (Crookes)

Scandium bromide, ScBi 611.0, is prepared in the same manner as the chloride. It forms rhombic crystals which at 120 are converted into 2ScBr, 31LO (Crookes).

Scandium perchlorate, Sc(ClO<sub>1</sub>), all O, from scandium hydroxide and perchloric acid, crystallises as a felt like mass of colourless, rhombic needles

Scandium iodate, Sc(10), 18H20, from scandium chloride and potassium iodate, is a white, crystalline powder almost insoluble in water, but soluble in nitric acid. When dired at 250' it becomes anhydrous, at intermediate temperatures, hydrates with 15H2O, 13H2O, and 10H2O may be obtained (Crookes).

## SCANDIUM AND THE OXYGEN GROUP,

Scandium sesquioxide  $\omega$  scandia,  $\mathrm{Se}_{2}\theta_{3}$  is obtained by igniting the hydroxide, carbonate, ovalate, nitrate, sulphate, etc., of scandium. It is a white powder of density 3 864, and specific heat 0 1530 (0° to 100°) 1 It is diamagnetic, the magnetic susceptibility being - 0.05 10 ° cgs, electromagnetic units per unit mass.<sup>2</sup> It dissolves slowly in cold, more readily in-

Nilson and Pettersson, Compt. rend., 1880, 91, 232. Ber., 1880, 13, 1459.
 R. J. Meyer and Wuotmen, Zeitsch. anorg. Chem., 1913, 80, 7

hot dilute acids, and very readily in hot concentrated acids, scandium salts and water being produced

Scandium hydroxide,  $\mathrm{Se}(\mathrm{OH})_{\mathfrak{p}}$  is obtained as a white, bulky, gelatinous precipitate by adding ammonium or sodium hydroxide to a solution of a scandium salt. When dired in the air it has a composition corresponding to the above formula. The hadrovide is a rather weak base, with a marked tendency to form basic salts. It readily dissolves in acids, forming salts which have a sweet, astringent taste

Scandium sulphide, Sc.8, is a yellow solid obtained by heating scandium sulphate in hydrogen sulphide. It is stable in an even at 100 but is decomposed by dilute acids or boiling water with evolution of hydrogen sulphide!

Scandium sulphite Sc (SO ), is obtained as a white procipitate by mixing solutions of scandium chloride and sodium sulphite (Crookes), by passing sulphur dioxide through an aqueous suspension of scandium hydroxide, the hexahydrate Sc<sub>2</sub>(SO<sub>4</sub>), 611<sub>2</sub>O, may be obtained. It is soluble in aqueous ammonium sulphite, and by evaporating the solution in an atmosphere of sulphin dioxide, seardium autorium entphile, Sc.(SO) (NII) SO 7H,O is obtained (Meyer and Diapier)

Scandium basic thiosulphate, SciOH/S,O<sub>n</sub> is obtained as a white, flaky precipitate when sodium thiosulphate is added to hot aqueous scandium chloride. Thorium and zircomum are smitully precipitated, but not the rare earth elements (Meyer and Drapier)

Scandium sulphate, Sc.(SO<sub>1</sub>), is obtained by disolving the oxide, hydroxide, or carbonate in sulphuric acid, and gently hertner until water and excess of sulphuric acid are climinated. It is a white powder of density 2-579, and specific neat 0-1639 between 0" and 100-2. The sulphate dissolves readily in water, at 12 there are 44.5 parts of anhydrous sulphate in 100 of the saturated solution (Crookes) - Unlike the rare earth sulphates, the solubility does not duminsh with rise of temperature - Scandium sulphate is not deliquescent and is insoluble in alcohol-

From a concentrated aqueous solution the hexabydrate, Se (SO<sub>4</sub>) 6H<sub>0</sub>O<sub>5</sub> crystallises out in small globular aggregates. This hydrate effloresees in dry an and leaves the pentahydrate Sc (SO<sub>1</sub>) 5H<sub>2</sub>O. The pentahydrate is the stable phase in contact with the solution at 25°, at which temperature 100 grains of solution contain 28.5 grains of anhydrous sulphate. The solubility alters with the addition of sulphuric acid, as shown by the following data -

| Grains of H SO, per litre                                     | . 00  | 24.5  | 19 0  | 121.5 | 213.3 |
|---------------------------------------------------------------|-------|-------|-------|-------|-------|
| Normality of H SO,                                            | 0.0   | 0.5   | 10    | 1.86  | 9 73  |
| Grams Se <sub>2</sub> (SO <sub>1</sub> ), per 100 of solution | 28 52 | 29.29 | 19.87 | 8 36  | 1.32  |

When dried over sulphune and the pentahydrate changes into the tetrahydrate, and this at 100° becomes converted into the dihydrate. The latter may be dehydrated at 250°

Scandium sulphate solution is only slowly and incompletely precipitated by oxahe acid and by sodium thiosulphate. Moreover, the equivalent

Wirth, Zeitsch anorg Chem., 1914, 87, 5
 Nilson and Pettersson, loc. cit
 Wirth, Zeitsch. anorg. Chem., 1914, 87, 9.

conductivity is abnormal, in that  $(\lambda_{10})_1 - \lambda_{12}$  is much smaller than would be anticipated. This is shown by the following data —

| Temperature, 25° C.                                |                |       |              |              |       |               |                                   |  |
|----------------------------------------------------|----------------|-------|--------------|--------------|-------|---------------|-----------------------------------|--|
| r =                                                | 32             | 61    | 128          | 256          | 512   | 1024          | λ <sub>1024</sub> λ <sub>12</sub> |  |
| Sc. 80()                                           | 25.3           | 33.8  | 10 1         | 17 7         | 57.6  | 72.1          | 13 8                              |  |
| ( (/80 <sub>i</sub> );<br>( Al, 80 <sub>i</sub> ); | 35 I<br>  51 I | 420   | 19 9<br>71 2 | 60 2<br>83 1 | 727   | 87 1<br>107 2 | 52 0<br>56 1                      |  |
| [Ci/s0]                                            | 67.1           | 7 × 3 | 90 7         |              | 119 ) |               | 60 7                              |  |

The explanation of these anomalous results is that scandium sulphate is really the scandium salt of a complex scandium sulphuric acid,  $H\left[\operatorname{Sc}(SO_4)_3\right]$ , thus,  $\operatorname{Sc}\left[\operatorname{Sc}(SO_4)_3\right] = \ln$  confirmation of this view it is found that whereas in migration experiments with scandium intrate and chloride solutions nothing abnormal is observed, with scandium sulphate a considerable quantity of scandium inigrates to the anode (Meyer and Bodlander)

A basic sulphate,  $8c_1O_1(28O_p)$  is produced when scandium sulphate is heated to dull reduces (Crookes). When scandium sulphate pentally drate is dissolved in sulphane and of density 1.6 acid scandium sulphate (or scandium sulphane and),  $8c_2(8O_1)_1(311_28O_1)_2$  (or  $11_2^28C_1(O_1)_3$ ), crystallises from the solution  $1_2^2$ .

Scandium potassium sulphate,  $Se_3(SO_4)$  3K  $SO_4$  or K  $Se_4(SO_4)$ , is obtained by adding an excess of potassium sulphate to a solution of scandium sulphate  $^2$ . It is moderately soluble in water and dilute potassium sulphate solution, but practically insoluble in a saturated solution of potassium sulphate. In this respect scandium resembles the metals of the cerite earths

Scandium ammonium sulphate, Sc<sub>3</sub>(SO<sub>1</sub>), 3(NH<sub>1</sub>)<sub>2</sub>SO<sub>1</sub>, resembles tho potassium salt closely (Meyer and Winter). When its aqueous solution is boiled, the sparingly soluble double salt, Sc<sub>2</sub>(SO<sub>1</sub>), 2(NH<sub>1</sub>)<sub>2</sub>SO<sub>3</sub>, separates out (Meyer and Diapter).

Scandium sodium sulphate, Sc<sub>2</sub>(SO<sub>1</sub>), 3Na<sub>2</sub>SO<sub>4</sub> 10H<sub>2</sub>O<sub>5</sub> can only be prepared in the presence of a large excess of sulphuric acid. It is readily soluble in water and in concentrated sodium sulphate, thereby resembling thorium sodium sulphate (Meyer and Drapier)

Scandium selenite, Sc<sub>3</sub>(SeO<sub>3</sub>), is an amorphous, white solid, insoluble in water. When heated with excess of aqueous selemous acid it is converted into a crystalline, insoluble acid scandium selenite. Sc<sub>2</sub>(SeO<sub>3</sub>), 311, SeO<sub>4</sub> (Nilson).

Scandium selenate,  $Sc_i(SeO_i)_0$ , crystallises from water in rhombic prisms of the octahydrate,  $Sc_i(SeO_i)_0$  8H O When dried at  $100^\circ$  the dihydrate is formed, and this is completely dehydrated at  $400^\circ$  (Crookes).

# SCANDIUM AND THE NITROGEN GROUP

Scandium nitrate,  $Sc(NO_4)_3$ , crystallises from a neutral aqueous solution as colourless, deliques cent, prisimatic crystals of the tetrahydrate,  $Sc(NO_3)_3$  1H<sub>2</sub>O,

With, Zeitsch. anorg. (hem., 1914, 87, 1 Nilson, loc. cit., Crookes, loc. cit.

<sup>2</sup> Nilson, loc cit.

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soluble in alcohol—The tetrally drate becomes pasty and anhydrous at 100°. Basic salts of the formulæ  $\mathrm{Se_2O_3}\,2\mathrm{N_2O_3}\,3\mathrm{H_2O}$  and  $\mathrm{Se_2O_3}\,2\mathrm{N_2O_3}$  have also been described (Crookes)

SCANDIUM.

The intrate is reachly decomposed by heat with the production of basic scandium intrates, and finally of scandia.

#### SCANDICM AND THE CARBON GROUP.

Scandium carbonate, Sc (CO)<sub>1</sub> 1211.0, is obtained as a white, bulky precipitate when a soluble carbonate is added to a scandium saft. It has the above composition when air dried. Scandium carbonate is rather inistable, and when dried at 100° it loses a little curbon drovide (Crookes).

Scandium carbonate dissolves in hot aqueous sodium or animonium carbonate. When these solutions are boiled, difficultly soluble scandium sodium carbonate, Sc $(\mathrm{CO}_{\mathrm{J}})$  (Na), O off O, and scandium ammonium carbonate, 28c (CO), (NH), CO offI.O, are obtained as crystalline precipitotes. These double carbonates dissolve without decomposition in a large volume of cold water, but when the solutions are boiled, basic salts or hydroxide separate (R J. Meyer and Winter)

Scandium oxalate, 8c (CQ<sub>1</sub>) 5H O, may be precipitated from an aqueous solution of a scandium saft by oxalic acid, and when an dried, forms a white, crystalline powder of the above composition. It is slightly, but distinctly soluble in water. In dilute in ls it is less soluble than the oxalates of the rule carth elements, as will be seen from figs. 19 and 21, and, unlike the rule earth oxidates, it is more soluble in dilute sulphinic than in hydrochloric acid. In these respects it resembles thorum oxalite, The solubility data are as follows (see fig. 19).

| Normality<br>of Acid | Hydro            | chlore | And              | Sulpl            | mpi | Acid   |
|----------------------|------------------|--------|------------------|------------------|-----|--------|
|                      | 25               | !      | 20,              | 25               |     | 50     |
| 0.1                  | 0.0209           |        | 0 (420           | 0.0355           | 1   | 0 0562 |
| 0.5                  | 0 0650           |        | 0.0870           | 0.0997           |     | 0 14*1 |
| 10 4                 | 0 1020<br>0 1715 |        | 0 1435<br>0 2 56 | 0 1663<br>0 3176 |     | 0 2493 |
| 50                   | 0 1170           |        | 0.655            | 0.7761           | i   | 1 1280 |

Boiling 10 per cent hydrochloric acid dissolves 0.8 per cent of its weight of hydrated scandium oxalate, the 20 per cent acid, 1.8 per cent of oxalate (Meyer and Winter)

(Meyer and Winter)
When dired over sulphune acid, the trihydrate, Sc (C<sub>2</sub>O<sub>4</sub>), 3H<sub>2</sub>O<sub>5</sub> is obtained, at 100° the dihydrate, and at 110° the monohydrate is produced (Crookes)

<sup>1</sup> R. J. Meyer and Wassjuchnow, Zeitsch unorg. Chem., 1914, 86, 284, of Wirth, abd., 1914, 87, 11.

Scandoum oxalate dissolves readily in hot ammonium oxalate solution, less readily in potassium or sodium oxalate, and on cooling, crystalline double oxalates of the type  $Se_2(C_2O_1)_1$   $3R_2^{\dagger}C_2O_1$   $10H_2O_2$  or  $R_1^{\dagger}[Se(C_2O_1)_1]$   $5H_2O_2$  separate out 1 Scandium oxalate has a strong tendency towards double salt formation,

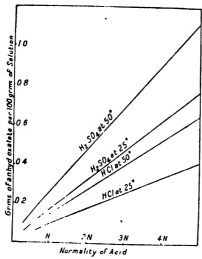


Fig. 19, - Solubility of scandium oxalate in acids

and when a solution of sodium or potassium oxalate is added to one of scandium nitrate. the crystalline precipitate first produced is a double oxalate of the type Sc2(C2O1), R'C2O4 (Meyer and Wassjuchnow)

Scandium acetylacetonate [(CII, (O),CH],Se, 18 obtained by adding scandium chloride solution to acetylacetone dissolved in a slight excess of ammonia - It may be purified by crystallisation from 70 per cent alcohol, or by dissolving it in benzene and precipitating with light petioleum By the former method it crystallises in colourless sparkling prisms; by the latter, in needles It is soluble in other and chloroform, and melts at 188° C, it crystallises in the rhombic

system and is isomorphous with the indium salt.2 Molecular weight determinations have been made by the freezing-point method in benzene, and by the boiling point method in benzene, chlorotorm and carbon disulphide, and the results show clearly that the molecular formula is that given above, and not [(CH, CO),CH],Se or [(CH, CO),CH],Se, with Se 29.

Under a pressure of 8 to 10 mm, scandium acetylacetonate begins to sublime at 157°, and sublimes rapidly at 187° without decomposition. At atmospheric pressure it melts at 188°, and commences to volatilise appreciably at 190°, no decomposition occurs below 250° 1. The acetylacetonate does not combine with annuous. In its properties it therefore resembles thorium acetylacetonate closely, and differs appreciably from the acetylacetonates of the rare carth elements

Scandium Salts of Organic Acids - Many of these salts have been described by Crookes

Scandium silicate (see p. 205)

<sup>• 1</sup> Witth, Zeitsch, among Chem., 1914, 87, 1 Meyer and Wassjuchnow (Loc ett.) describe the salt Sc<sub>2</sub>(C<sub>1</sub>(1)<sub>4</sub>)<sub>7</sub> 4(NH<sub>4</sub>)<sub>2</sub>(C<sub>2</sub>O<sub>4</sub> 7H<sub>2</sub>O)

• Jaeger, Proc K. Akad Wetensch Amsterdam, 1914, 16, 1095, Rec trav. chim., 1914,

<sup>33, 342</sup>Meyer and Winter, loc, cit. Meyer, Zeilsch. anorg. Chem., 1914, 86, 288; Morgan and Moss, Trans (them. Soc., 1914, 105, 196.

Morgan and Moss, loc cit

### SCANDIUM AND BORON.

Scandium orthoborate, 8clio, prepared by fusing bone anhydride with scandia and extracting the excess of bone anhydride with water, is a white, insoluble powder (Crookes)

### DELLCHON AND ESTIMATION OF SCANDIUM.

Minute amounts of scandami may be detected in immerals, etc., owing to the extreme dehency of its spark and are spectra. When utilising the spirik spectrum it is first necessary to decompose the substance under examination, precipitate the carths as oxalites, igiote, and dissolve the residual oxides in hydrocletoric acid. Owing to the slight solubility of scandium ovalete in hydrochloric acid, the filtrate from the oxalates should be precipitated with ammonia, and the precipitate likewise tested for se indimit. In examining rare cuth immerals it is, moreover, necessary to concentrate the scandina into a small fraction of the rare earths. It is also necessary to carry out a prolumnery chemical treatment before exagining the are spect a when it is earth inmends are being examined, but with other numerals and rocks it suffices to powder the substance, heat it to redness in order to shave off water and gises, and then examine the arc spectrum or the powder in the usual way, using a carbon are. At least 0.5 gram must be used, and it must be completely vaporised if traces of scandium are to be detected ! The spectroscopic investigation should be made by the photographic method, the region from 3500 to 3700 being particularly exammed (Crockes)

The most intense lines in the are and spark spectra of scandium are as follows

```
spark | 3353 89, 3558 71, 3567 88, 3572 72, 3576 53, 3581 15, 3613 98,
       3630'90, 3642'96, 3651'96, 4247'00, 1311'25, 4320'91, 1325'15, 137'16', 1100'50, 1115'72 1353'89, 3372'33, 3558'69, 3567'88, 3572'72, 3576'53, 3613'98, 3630'90, 3612'96, 3907'69, 3912'03, 4020'55, 4023'83, 4217'00,
        4314 25, 4320 91, 4325 15, 4374 68, 4400 56, 4415 72, 6305 88.
```

The are spectrum of scandium consists of two distinct sets of lines which behave very differently in solar spectra. One of these sets corresponds to the "enhanced" lines of other elements. The are spectrum when taken in air shows very characteristic flutings alue to scandium oxide. They disappear when the are is taken in hydrogen, and at the same time the spectrum ; approximates to that observed with the spark discharge (Fowler)

For the quantitative estimation of scandium it is precipitated as the oxalate and weighed as the sesquioxide

Eberhard, Sitzungsber K. Abad. Wiss. Berlin. 1908, p. 851. 1910, p. 404. Chem.
 Neus, 1909, 99. 30, 1910, 102, 211.
 Lockyer and Baxandall, Proc. Roy. Soc., 1905, 74, 538. Fowler. Phil. Trans., 1908.
 A. 209, 17. Proc. Roy. Soc., 1908. A, 81. 335. Finer and Hassleck, Die Spektren der Llemente bei nurvalient Iruck (Lepzig and Virina, 1911), vol. 1., Kayser, Handbuch der Stektenburg (Lowing, 1912), vol. 43. Spektroskopie (Leipzig, 1912), vol vi p 144.



# CHAPTER X.

### THE RARE EARTH ELEMENTS.

Introductory.—"Chemists distinguish such substances by the name of pure earth, as are brittle, incombustible, infusible by the heat of furnaces, not soluble in several hundred times their weight of water, and destitute of metallic splendom. There are few carthy substances which may not be reduced by analysis to one of the five following primitive earths -the siliceous, argillaceous, calcareous, ponderous, and magnesian carths or otherwise, taken substantively, they are called siley, clay, line, barytes, and magnesia "1

The preceding quotation will serve to show the meaning attached by chemists to the term earth at the time when the foundations of modern chemistry were being laid. Five earths were recognised, viz., silica, alumina, baryta, lime, and magnesia, and the existence of another (strontia) strongly suspected. Glucina or beryllia was added to the list in 1798, and a number of further additions were made within the next twenty years from that date It will be noticed that the earths all agree in one respect, viz, they are exides which can only be reduced with great difficulty. That the earths were oxides was suspected by Lavoisier, and Davy's classic experiments (1808) on the electrolysis of the alkalies and alkaline earths afforded the first experimental confirmation of this view. By Berzehus, Guielin, and others the earths were usually divided into two groups, (i.) the alkaline earths (hine, strontia, baryta, and magnesia), and (ii) the earths proper, their metallic constituents were classified, together with the alkali metals, as light metals. Moreover, silica was sometimes removed from the list of earths and placed among the acids

For a considerable number of years it has been customary to refer to cortain earths as the rare carths, since for many years after their discovery the only available sources of them were a number of rare minerals, found principally in Scandinavia. Originally, two such earths were recognised, ceria and yttma, these untual rare earths, however, have proved to be so extremely complex that at the present time fifteen rare earths receive official recognition, and it is probable that the existence of one or two more will be substantiated.

There is no precise limitation to the nature of the earths that are to be regarded as rate earths. In this book the term rate earths is restricted to the sesquioxides,  $M_aO_3$ , that have been recognised as components of the original "cena" and "yttim" and to corrum sesquioxide. Accordingly the earths gluena, thora, and zircoma, and the earth-acids, viz, titame oxide and the pentoxides of columbium and tantaliin, are not discussed, although

some authors include them among the rare earths. Moreover, scandia is not included among the pare earths, but is given separate treatment (Chapter IN).

The rare earth elements, their symbols, and atomic weights are given in the following table in which the elements are divided into three groups for reasons that will be subsequently explained (p. 219)  $\rightarrow$ 

| Cerman                                                       | Cerum Group Terbinin Group |                                             |                                  | Yttimin Group    |                         |                                                                                               |                                             |                                                    |
|--------------------------------------------------------------|----------------------------|---------------------------------------------|----------------------------------|------------------|-------------------------|-----------------------------------------------------------------------------------------------|---------------------------------------------|----------------------------------------------------|
| Lanthanum<br>Ceruum<br>Prascodyumum<br>Neodyumum<br>Samarium | La<br>Pr<br>Nd<br>Sm       | 130 p<br>110 2 ;<br>140 0<br>141 ;<br>150 4 | I mopuum<br>Gadohurum<br>Terbium | Fu<br>tol<br>116 | 152 0<br>157 3<br>159 2 | Dysprosium<br>Helimium<br>Vitrium<br>Erbanin<br>Thalium<br>Viterbanin<br>Lateerium<br>Celtuum | Dy<br>Ho<br>Y<br>En<br>Im<br>Yb<br>Lu<br>Ct | 162 5<br>163 5<br>85 7<br>167 7<br>168 5<br>17 1 5 |

From the analytical point of view, the rare earth elements resemble aluminium and chromium, from other points of view they resemble the alkaline earth metals and bismuth. Their most striking characteristic, however, is their extraordinary chemical similarity one with another, and since the rare earths are always associated together in nature, then separation from one another is a very formulable task. The difficulties encountered far surpass those met with in separating, say, calcium from strontium, or mickel from cobalt, in the separation of the rare earth elements, analytical methods similar to those ordinarily employed in analysis are restricted to the separa tion of cerum from the others

In the present chapter a general account of the chemistry of the rare earth elements is given. In Chapter XI the separation of the rare earths is discussed, and in Chapters XII, XIII, and XIV, the individual rare earth elements and their compounds are described

Occurrence — In minute quantities, the pare earths are extremely widely disseminated in nature. This has been shown especially by experiments dealing with cathodic phosphorescence spectra \*\*\* Traces of rare earths occur in many immerals, of which scheelite," 5,6 pitchblende, apartite," 6 certain varieties of calcite, "fluorspin, " assitente," and woltramite may be particularly mentioned. Bare carths have also been detected in coral, " bones,2,6 the ashes of rice and tobacco, 6 etc.

The number of immeral species containing the rare earths is very large They consist for the most part of silicites, tit mates, zirconates, tantalates,

<sup>&</sup>lt;sup>1</sup> See Schilling, Das Vorkommen der seleen Lehn im Menerativehe (Munich and Berlin, 1904), Brogger Zeilich Kryst Min, 1800–16, und Die Menerative andnorwegischen Grand Permatitionige, 1904, Chen and Westten, The Meneration of the Ruser Elements (\*\*Griffin & Co., 144, 1912), Levy, The Kare Virths (Arnold, 1915), and the larger mineralogical text books of Dana, Hintze Docker, etc. <sup>2</sup> Grookes, Phil Trans., 1883, 174, in. <sup>801</sup>. Comm. News, 1884, 49, 159, 169, 181, 194, 205.
<sup>3</sup> Urbain and Scal, Compt. rend., 1907, 144, 20.

<sup>1, 206

\*\*</sup>Uthain and Scal, Compt. rend., 1907, 144, 20 \*\*

\*\*Uthain, Ann. Chim. Thys., 1909, [vin.], 18, 356

\*\*de Rohden, Compt. rend., 1914, 159, 318., Ann. Chim., 1915, [ix.], 3, 338.

\*\*Cossa, Guzzelta, 1879, 9, 118.

\*\*Ossa, Guzzelta, 1879, 9, 118.

\*\*Headden, Amer. J. Nr., 1906, [iv.], 21, 391.

\*\*Humphreys, Astrophys. J., 1904, 20, 266

\*\*Eberhard, Sitzungsber. K. Akad. Wiss. Berlin, 1908, p. 851, 1910, p. 404

and phosphates, in which the rare earths are associated with lime, theria, glucina, oxides of iron, uranium, etc. They occur principally as included minerals in granite and pegmatite.

The minerals occur mainly in four or five localities. The Scandinavian deposits were the first known, and, although not very plentful, are extremely rich in mineral species. In Norway, the chief localities are near Langesund Flord, and include Arendal, Brewik, Kragero, Hittero, Risor, and Moss, the Swedish deposits are found at Ytterby and Bastinas. The graintes and pegmatites of the coast of Greenland are also rich in these minerals, which are almost invariably associated with arguite. In North America, deposits occur in North and South Carolina, Colorado, Vriginia, Idaho, Maine, and particularly at Barringer Hill, Danio Co., Texas I. In South America, largo deposits of monazite sand occur in Brazil, particularly in the previnces of Bahia, Minas Geraes, Espirito Santo, Matto Grosso, and Goyaz. Other deposits of rare earth minerals are found in the Urals, particularly at Miask.

Among the most important time earth innerals are cerite, allunite, monazite, eschynite, gadolinite, senotime, fergusonite, ythrotantalite, simurshite, polycrase, and encents. The first four of these numerals are rich in ceria earths and poor in yttra earths, the reverse being true of the others.

Cente consists essentially of a hydrated scheate of ceria, with small amounts of iron, calcium, etc. It crystallises in orthorhombic prisms (a b c -0.999 1 0.813), but commonly occurs in the massive or granular form, and contains 59 to 72 per cent of ceria. The amount of yttrac present may reach 7 per cent. Hardness, 5.5., density, 1.86 to 1.91. The colour varies from red to brown or grey. Cente is found embedded in gnoise at Bustnes, associated with mich, horibblende, all unite, and chalcopyrite.

Allante or orthite (varieties, bucklandile, uralorthite, bagrationite, standium-orthite, muromontite, bodenite) is a basic orthosilicate of calcium, aluminium, iron, and cerimi. It crystallises in monochine tables, plates, or needles (a b c=1551-1 1769,  $\beta$ -64°59') of a reddish black colour, the crystals often resembling rusty ualls, and generally contains from 16 to 25 per cent of rare earths. Hardness, 5.5 to 6, density, 3.5 to 4.2. Allante is found in Texas, Greenland, Norway, and Sweden

Monards, Ce(La,D)PO<sub>4</sub>, is essentially cerum (lanthanum, etc.) orthophosphate. It forms monodime crystals (a. b. 0.969–1.0.926,  $\beta$  = 76°20′) which vary in colour from hyaemth red to brown. Hardness, 5 to 5·5, density, 5.0 to 5·2, and occasionally as high as 5.5. It generally contains 60 to 70 per cent of ceria, and from traces to upwards of 5 per cent of vitua

Monazite crystals, which are rather scarce, are found in granites and granite gnesses in Norway and Greenland, in Ceylon, and in one of two other localities. Rolled grains of monazite, however, occur in abundance, being found in large deposits of said and gravel in various localities. These monazite saids have been produced by the weathering of rocks which originally contained a very small percentage of monazite, and the subsequent washing away of the lighter materials produced. The monazite is associated with numerous other numerals, e.g. quarte, thorite, zucon, garnet, tournaline, hornblende, augite, obvine, topaz, spinel, rutile, magnetite, titainte and easiterite.

Hidden and Mackintosh, .tmcr. J. Sci., 1889, [iii.], 38, 474, 1905, [iv.], 19, 425
 By "ceria" and "yttria" throughout this section are meant "earths of the cerium group" and "earths of the yttrium group" respectively.

In North and South Carolina monazite occurs in stream beds and placer deposits, its origin being the granitic biotite-gners, and dioritic hornblendegness of the surrounding country. The gravel deposit varies from one to three feet in thickness The Brazilian deposits occur in the provinces already mentioned (p 218) They occur along the coast and also mland, following the course of certain water-traversed syemite rocks and gnesses. Monazite sand is also found in India in the native State of Travancore, im Ceylon, in various localities in the Federated Malay States, and the protected States of Kedah and Kelantan,2 and in various parts of Nigeria 3

Monazite sand is of considerable commercial importance, since all monazite contains a little thorum, which is employed in the manufacture of in-candescent gas mantles. The commercial value of a sample of monazite sand depends, in fact, upon its thorum content

Most of the world's supply of monazite sand has come from Brazil, where the annual production of sand containing 90 per cent of monazite exceeds 6000 metric tons. Monarite separated from the concentrated sands found on the sea coast contains 5 to 7 per cent of thoria, monazite from the inland deposits contains 1.0 to 5.7 per cent of thoria. Prior to its exportation, the sand and gravel is washed in sluce boxes like placer gold, and the dried sand freed from magnetite, etc., by an electromagnetic process. It is not difficult to "concentrate" the sand in this way till it contains 95 per cent of monazite. The thoria content of the product regularly marketed varies from 5.75 to 7.1 per cent. The monazite's ind deposits in Carolina are poor in thoria and of little or no present commercial value, but there is a very considerable production of the sand, very rich in thoria, from Travancore

The composition of a number of simples of monasite is given in the following table -6

| . <del></del> · · <del>-</del> -                                                                                                                    | Espuit Sant                                        | Ve ban<br>Buzil                                 | North Carolina         | R. sapara,                                              | Travancon<br>Exch<br>North Nigeria | Flare R .et S. ath N.gerea              | Pehnig Malay                                   | Kalaam,<br>State of Kedan                     |
|-----------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------|-------------------------------------------------|------------------------|---------------------------------------------------------|------------------------------------|-----------------------------------------|------------------------------------------------|-----------------------------------------------|
| ThO,<br>C(t)<br>In,O <sub>e</sub> ct<br>Y <sub>1</sub> O <sub>e</sub> ct<br>Fe <sub>2</sub> O <sub>e</sub><br>Al <sub>2</sub> O <sub>e</sub><br>CaO | 6 06<br>1 6 2 12<br>1 0 50<br>0 07<br>0 10<br>0 21 | 6 50<br>61 40 {<br>0 70<br>1 50<br>0 08<br>0 30 | 6 19<br>31 3-<br>30 88 | 10 29<br>27 37<br>30 13<br>2 11<br>0 81<br>0 17<br>0 41 | 1 50 3 00<br>0 17   0 35           | 31 40<br>29 20<br>2 09<br>0 75<br>0 0 5 | 25 16<br>32 72<br>2 80<br>0 84<br>2 78<br>0 61 | 3 53<br>64.05<br>2 40<br>0 64<br>0 07<br>0 17 |
| S <sub>1</sub> O <sub>2</sub> .<br>P <sub>2</sub> O <sub>5</sub> .<br>Loss on ignition                                                              | 0.75<br>28.50<br>0.35                              | 0 64<br>28 46<br>0 64                           | 1 40<br>29 28<br>0 20  | 1 03<br>27 67<br>0 20                                   | 9 90   1 20<br>26 82 26 29         | 0.82                                    | 0 92<br>23 92<br>1 28                          | 1 08<br>27 87<br>0 52                         |

Tupper, Rev. Geol. Survey, India, 1914, 44, 186., Ch. at Comm. J., 1915, 34, 205.
 Bull. Imp. Inst., 1906, 4, 391., 1911, 9, 39
 See S. Johnstone, J. Sov. (Iran. Ind., 1914, 33; 55.

See S. Johnstone, J. Nov. Chem. Line, 1914, 33; 50.
 Mining Industry, 1909, 18: 557
 Gottschalk, Chem. Eng., 1915, 21, 169., Mining Engineering World, 1915, 42, 103
 Data taken from a paper by S. Johnstone, J. Nov. Chem. Line, 1911, 33: 55.
 See also Genth, Amer. J. Not., 1889, 38, 203. Blomstand, J. pinkt Chem., 1890, [n], 41, 286; Chem. Soc. Abis., 1890, 58, 111, 571.
 1891, 60, 1168, Glaser, J. Amer. Chem. Soc., 1896, 18, 782, Schilling, Zeitsch. angew. Chem., 1902, 15, 869

The preceding results refer to monazite carefully freed from admixed minerals.

. Eschente,  $R_2^m(Cb_1O_{12},R_2^m(T)Th)_2O_{19}$  is a columbo-titanate of cerum , it contains a considerable quantity of thornin. It crystallises in orthorhombic prisms of tables (a-b-c=0.487-1-0.674), which are usually black in colour—Hardness, 5 to 6, density, 4.93 to 5.17—Eschenite contains 15 to 24 per cent of cena, and 1 to 3 per cent of yttra. It is found in Norway and the Utals

Gadolinete is a basic orthosiheato of yttimm, glucmum, and iron forms monoclinic prisms (a b c 0.6273, 1.13215,  $\beta$  89° 26.5'), and occurs in large masses, which are dark in colour and give a greenish grey streak. Hardness, 6.5 to 7, density, 4.36 to 4.47. It contains 36 to 48 per cent of yttria, and 5 to 16 per cent of ceria. Gadolimite is found as crystals in granitic pegmatite in Norway and Sweden, and occurs abundantly in Llano Co., Texas 2

Xenotime is essentially yttimm orthophosphate, YPO<sub>4</sub>—It forms tetragonal crystals (a |c|=1 0.6187) resembling zeron. Hardness, 1 to 5, density, 4.15 to 156 It contains 51 to 61 per cent of yttim, and from 8 to 11 per cent of ceria. Aenotime occurs in guerss, granites, syemites, and in their pegmatite veins in Norway, Sweden, Switzerland, it is also found in the monazite saids of Carolina, Georgia, and Brazil

Fergusonite, (Y,Ce)(Cb,Ta)Op is a metatautalate and columbate of yttimm It crystallises in tetragonal bipyramids (a c 1 1 161) of brownish black coloni, and gives a pale brown streak. Hardness, 5.5 to 6 , density, 5.8 . It contains 30 to 16 per cent of vttria, and upwards of 9 per cent of ceria, also thorum and manium in considerable quantity. Fergusonite was discovered in quartz in Greenland), it occurs in Texas, Virginia, and Carolina, associated with samarskite and gadolimite, and in Norway and Sweden, associated with envenite

Pttrotantalite and samar-kite are columbo tantalates of yttimm, iron, and calcium. Samarskite differs from yttrot utalite in being rich in uranium Both innicials are orthorhombic and reddish-brown to black in colour Hardness, 5 to 6, density, 5.5 to 5.9. Attrotantalite contains 17 to 38 per cent of yttra, and upwards of 2 per cent of cera, samarskite, 5 to 21 per cent of yttria, and from 2 to 5 per cent of ceria. Both numerals occur in granite pegmatite in various localities, samarskite occurs massive in North Carolina in large quantities 3

Eurenite is a columbo-titanate of yttrium, which contains uranium orthorhombic (a · b  $\epsilon$  = 0.364 / 1 (0.303), but usually occurs in brownish-black masses which have a pitchy lustic and give a yellow to reddish brown streak. Hardness, 6.5, density, 4.6 to 5.0. It contains 13 to 30 per cent of yttria, and 2 to 8 per cent of ceria Euxemite is found in Norway and Greenland.

Polycrase is similar in composition to envenite, with which it is iso-

<sup>1</sup> For a bibliography of the literature on monarite so hat as 1894, see Nitze, Settenth Ann Rep. U.S. God Narry, 1894-95, p. 667. For discussions of the constitution of monarite, see Dumington Amer. Chem. J. 1882, 4, 138, Penfield, Amer. J. See, 1882, 24, 250, 1888, 36, 322; Ramnelsberg, Zeitsch Krust. Min., 1879, 3, 101. Blomstrand, loc. etc., and Zeitsch. Krust. Min., 1887, 6, 160, 1892, 10, 367, Kress and Metyger, J. Imer. Chem. Soc., 1909, 31, 610. Johnstone, loc. etc., Chem. Soc., 1909, 31, 610, Johnstone, loc. etc., Genth, Amer. J. See., 1889, [in.], 38, 198, see also Bull. U.S. Geol. Survey, 1908, No. 340, 286.

3 Bull. U.S. Geol. Survey, 1891, No. 74

morphous, but contains less columbium and more titanium. The present day opinion is that cuvenite and polyerase are members of an isomorphous series of mixed yttrium metacolumbate and inctatitanate, and the name cuvenite is retained for mixtures in which the ratio Cb O<sub>2</sub>. TrO<sub>2</sub> exceeds 1 - 2, while the name polycrase is given to the others. Euvenite is dimorphous, the less common crystalline form being called priorite. Polyerase is likewise dimorphous, the other modification being known as blomstrandine. Priorite and blomstrandine are isomorphous minerals Thus the cuvente group of minerals constitutes an isodimorphous series 1

Among the other rare earth minerals may be mentioned tysoute, condylite, a fluorarbonate of cermin and barnin, bastuasite, Ce,(CO), CeF, anoplete, 4Ce(OH)CO 38rCO 3H,O, lanthanete, La (CO) 8H O tertomete, a fluosilicate of cermin, thornin, and calcium, erdinamite, a mineral resembling allanite, beckelete, (Ce,Y),(SiZi),Ca,O1, hellandite, movambete, and rinkite, cerum silicates, tengerile,  $Y_2(CO_3)_3$ , cappelente, a yttimin borosilicate; rowlandite,  $X_1S_1O_{12}$ , yttrodite,  $Y_2S_2O_{12}$ , thatente,  $\Pi[X_1S_1O_{12}]$ , yttrocravite, a vttrium and thorium titunate, keithanite (or yttroditimite), an isomorphous mixture of titanite, Ca(TiSi) O, and (Y,Fe,Al) SiO, poinchlore, a columbate of cerum and calcium, etc., hoppite, 5Ce<sub>2</sub>Cb<sub>3</sub>O<sub>7</sub> 2Nal resorted a vittum columbate, YChO<sub>1</sub>, with an 1 omorphous admixture of yttrum metathanate  $Y_2(\text{TiO}_1)_1$ , separate YChO<sub>1</sub>, helmite and locarskite, tantilates of yttrum, etc.

The rare earth numerals have many points of interest, which cannot be discussed in this book 5. It may, however, be mentioned that the researches of Boltwood" and Strutt? have shown that in general the rare earth immerals are strongly radioactive, and that, with the exception of a tew manning numerals, scurcely any others exhibit more than a feeble radioactivity rule, therefore, the rare earth immerals are rich in occluded helium. The namerals owe their radioactivity to the presence of thorium or manium or both. The almost invariable association of these elements with the rare earths can scarcely be fortuitous, but no explanation of it is yet known. In some cases the thorum or manning is in excess and only small amounts of rare carths are present, e.g. thorounite and pitchblende, in other cases the reverse holds good, e.g. monutite. Indeed, in the cases of monazite and a number of other immerals the small amounts of uranium present were overlooked in analyses until with the development of the study of radioactivity their probable presence was inferred

Historical —(i) Early History -In 1794 the Funnsh chemist Gadolin isolated a new carth or oxide from a black immeral (gadolinite) found at

See Biogger, Vol. Sel. k. Skritler, Cher tearner, Math. Nature. Ed., 1906, No. 6, 1;
 Abst. Chem. Soc., 1907, 92. ii., 885., Lange, Zeesh. Nature es. Harb., 1910, 82., Abstr. Chem. Soc., 1911, 100, ii., 199
 T. Vogt, Centr. Min., 1911, p. 373., Inhib. Min. 1914, H. 9
 Hauser, Bir., 1907, 40, 3118. Zeitsch. among. Chem., 1908, 60. (20)
 Mallet, Amer. J. Ser., 1877, [iii.], 14, 397
 An interesting and faulty complete account of the properties of the commercials will be found in Levy, The Rare Eacths (Arnold, 1916)
 B. B. Boltwood, Amer. J. Ser., 18001, [iv.], 18, 97., 1905, [iv.], 20, 253., 1900, [iv.], 21, 415., 1908, [iv.], 25, 269., Phil. Mag., 1905, [iv.], 9, 599
 Strutt, Proc. Roy. Soc., 1904, 73, 191; 1905, A, 76, 88, 312., 1907. A, 80, 56, 1908, A, 80, 572

A, 80, 572
On the occurrence of helium in minerals see Vol 1, pt. u.

Ytterby, near Stockholm! This discovery was confirmed in 1797 by Ekeberg,2 who showed that Gadolin's "earth" was a mixture of the then recently discovered gluema and a new earth to which he gave the name of yttria. Mom this "yttria" Berzehus and Gahn extracted a little ceria in 1816.s

In 1804 another Swedish mineral, known as the "heavy stone from Bastnas," was evamined by Berzelius and Hisinger in Sweden, and by Klaproth in Germany. It was found to contain a new earth, which Klaproth called ochroite, because it turned dark yellow when heated. The Swedish chemists called it certa, i.e. oxide of cerium, and named the nineral certie.

In 1839 Mosander showed that "certa" was complex, for when cerous

hydroxide was suspended in potash and treated with excess of chlorine, part of it went into solution and the remainder was converted into a cition-yellow higher hydroxide Retaining the name cena for the earth corresponding to the yellow hydroxide, he named the earth that had passed into solution lanthana or oxide of lanthanum.6 Subsequently he found that an approximate separation of lanthana from ceria could be made by extracting crude "ceria" with 1 per cent mitric acid, the lanthana passing into solution. For various reasons, Mosander considered that lanthana was white, the brown colour of his preparations being due to foreign matter, and early in 1840 he obtained proof of this hypothesis, and succeeded in showing that a third earth existed in crude "ceila". This earth, which he found to be responsible for the brown colour of his ceria and lanthana and for the pale amethyst colour of his cerous and lanthanum salts, he named didymoa or oxide of didymium.7 Mosander found that the fractional crystallisation of the sulphates was the best method for separating lanthana and didymia

The complex nature of "yttina" was suspected by Scheener in 1812 s and proved by Mosander in 1813 " By the fractional precipitation of "yttinum" salts with ammonia, and with acid potassium oxalate, he resolved "yttim" into three oxides. The most basic oxide, which was present in largest amount, he called yttria or oude of yttrium, of the other two, one, which was pink, he named terbia or oxide of terbium, while the other, which formed a brown perovide, he named erbia or oxide of erbium basicity decreased in the order yttina, cibia, terbia.

Mosander's cerium, lanthanum, and yttrium are to-day classed among the chemical elements. It is now known, however, that his "didyminin," "erbium," and "terbium" were complex, in particular, that his perovide-forming element "erbium" contained only a trace of an element capable of

<sup>&</sup>lt;sup>1</sup> Gadolin, K. Svenska Vet.-Akad. Handl., 1794, p. 137; Crell's Annalen, 1796, 1-313.

Gadolm, K. Svenska Vet.-Akad. Handl., 1794, p. 137; Crell's Annalen, 1796, p. 1313.
 Ekcberg, Crell's Annalen, 1799, p. 433, K. Svenska Vet.-Akad. Handl., 1802, p. 68.
 Berzelnus and Galm, Schwerger's J. 1816, 16, 250, 404.
 Klapnoth, Sitzungsber K. Akad. Wiss. Berlin, 1804, p. 165
 Berzelnus and Hisinger, Gehlen's alla J. Chem., 1804, p. 165
 Berzelnus and Hisinger, Gehlen's alla J. Chem., 1804, 2, 303, 397, Ann Chim. Phys., 1804, 50, 140.
 The discovery was announced by Berzelnus. See Compt. rend., 1839, 8, 356, Pogg. Annalen, 1839, 46, 648.
 47, 207, Annalen, 1839, 25, 255, 503. For Mosander's own account of his work see Mosander, Phil. Maq., 1843, [iii.], 23, 241; Ann. Chim. Phys., 1844, [iii.], 11, 484; Pogg. Annalen, 1848, 60, 297, Annalen, 1843, 48, 210.
 Hee carly methods employed in the separation of the salts of cerium, lanthanum, and didymium, see Mosander, Loc. cit.; Marignac, Ann. Chim. Phys., 1849, [iii.], 27, 209, Watts, Quart. J. Chem. Soc., 1850, 2, 140.
 Scheerer, Pogg. Annalen, 1842, 56, 479.
 Mosander, Loc. cit.

forming a perovide by the ignition of its hydroxide, nitrate, or oxalate. Moreover his yttria was by no means pure

In 1857, shortly after the invention of the spectroscope, Gladstone observed the absorption spectrum of "didyimum," and in 1862 Bahr discovered the absorption spectrum of "terbnim". The application of the spectroscope to the investigation of the rare earths was extended by Delafontaine and others Meanwhile, doubt was cast on the existence of Mosander's "erbium." In 1860 Berlin introduced his classic method for the fractionation of the gadolinite earths, namely, the partial decomposition of the intrates by heat. He failed to obtain the peroxide-forming earth, and unfortunately gave the name of "crbia" to Mosander's pink earth This incorrect designation has been adopted by all subsequent workers.

The results obtained by Berlin were confirmed by Bahr and Bunsen in 1866, and by Cleve and Hoghand in 1872. According to these chemists, Mosander's "peroxide of erbium" was really a trace of peroxide of "didymium "4 The existence of Mosander's "erbium," however, was vigorously upheld by Delafontame."

In 1873 the rare earth mmeral samurskite was found in considerable quantity in a ninea mine situated in Mitchell County, North Carolina In 1877 comparatively large amounts of this immeral became available,6 and the chemistry of the rare earths entered upon a new phase. Almost the first result of the study of the samarskite carths was the definite establishment, by Delatontaine, of the existence of Mosander's "erbia," which ho was obliged to remaine "terbia" as the name "terbia" was by that time in common use for Mosander's "terbia" Almost simultaneously, Marignac showed conclusively the occurrence of Mosander's "erbia" in the gadolinite earths. The existence of the present day terbia was thus placed beyond question, its isolation in a state of purity, however, was not accomplished until nearly thirty years later 9

Whilst engaged in the analysis of samarskite, Lawrence Smith 10 observed indications of the existence of a new rare earth, and early in 1878 he aumounced the discovery of mosandrum, a new rate earth element resembling the elements of the cerite earths 11. The discovery was adversely criticised by Delafontaine and Marignac, who regarded "niosandrium" as identical with their "terbium," 12 and at no time was the existence of "mosandrum" generally admitted Eight years after the announcement of

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Glubstone, Quart J. Chem. Soc., 1858, 10, 219, 1859, 11, 36
 This fact is referred to in numerous early papers. Apparently, no description of the spectrum was published by Bahi until 1866 (Bahi and Bunsen, Annalen, 1866, 137, 1).
 Berlin, Forbandl. Skand. Nat. Kyob., 1860, p. 448.
 Bahi and Bunsen, Annalen, 1866, 137, 1., Cleve and Hoglund. Bull. Soc. chim., 1872, pp. 1-8, 1941-250.

<sup>Delafontaine, Arch. Sci. phys. and., 1864, 21, 97, 1865, 22, 30, 1866, 25, 105, 1874, 51, 48
See Lawrence Smith, Amer. J. Sci., 1877, [iii], 13, 359.
Delafontaine, Arch. Sci. phys. and., 1877, 59, 176., 1878, 61, 273. Ann. Chem. Phys., 1878, [v], 14, 248.
Maniguae, Arch. Sci. phys. and., 1875, 61, 283, Ann. Chim. Phys., 1878, [v], 14, 247.
Utbain, Compt. rend., 1905, 141, 521.
L. Smith, Amer. J. Sci., 1877, [iii], 13, 359.
L. Smith, Compt. rend., 1878, 87, 146, 148, 831, 1879, 89, 480, Amer. Chem. J., 1833, E. 44, 73.</sup> 

<sup>1883, 5, 44, 73

18</sup> Delafontaine, Compt rend , 1878, 87, 600; Mangnac, abid., 1878, 87, 261.

its discovery, Lecoq de Borsbaudran 1 showed the complexity of "mosandrum," and from his results it can now be stated that in addition to terbium there were present didymnum (a trace), samarium, dysprosium, and probably gadolimum in the so called element

Shortly after the announcement of the discovery of "mosandrini," Delaiontaine claimed to have isolated a new earth of the yttimm group, which he named oxide of phillipium 2. The history of this so called element may be conveniently given at this point, as subsequent work has shown "phillipium" to have been nothing but a complicated mixture. The salts of "phillipium" were said to be characterised by an absorption band \$\lambda 1490, an unfortunate method of describing the element, since erbium salts have an absorption band in the same position. Soret, who pointed this out, was inclined to regard "phillipium" as identical with an element  $\mathbf{X}$  which he had characterised just previously by a particular absorption spectrum.<sup>3</sup> Delafontame at first admitted the possibility that "phillipinin" was complex, though he thought it unlikely, but, almost immediately afterwards, he concluded that "phillipnim" had no absorption spectrum, attributing the band \$1490 and others he had observed to the presence of a little of Soret's X as an impurity 1 In 1882, however, Roscoe showed that "phillipium" was only a mixture of yttrium and the "terbium" of that period, and his conclusions were supported by Crookes? The later work of Urbain's has confirmed the view that "phillipium ' was only a mixture, though Delafontaine asserted its clementary character as late as 1897 ?

From this point in the history it is perhaps clearer to deal separately with the cerite and gadolinite earths

(u) The Cerum Group - About the same time that he announced the discovery of "phillipnim," Delafontaine stated that in exagining "didymia" extracted from samarskite, he had isolated a new earth, oxide of decipium, characterised by two bands in its absorption spectrum at \$\lambda 1160 and \$1780. The year following, Lecoq de Boisbaudian isolated a new earth from samarskite "didymia," and called it . imaria or oxide of samarium s The band \$1160 was also present in the spectrum of this element, and samanum would therefore appear to be more properly called decipium but for the fact that, a year or two later, Delafontaine 10 declared his "decipia" to be a mixture of two earths, one being samaria and the other an earth the salts of which gave no absorption spectra. He reserved the name oxide of decipium for this second earth. It is almost certain that this is to be identified with the earth Ya discovered in the meantime by Marignac Delafontaine's designation for the earth must therefore give way to Mangnac's, and so the name "decipium" disappears from the list of rare earth metals

Lecoq de Barbandran, Compt. rend., 1886, 102, 647.
 Delafontame, Arch. See phys. nat., 1878, 61, 273. Compt. rend., 1878, 87, 559
 Soret's element A is discussed later (p. 225)
 Soret, Compt. rend., 1879, 89, 521, Delafontsme, ibid., 1880, 90, 221, Arch. See. phys. nat., 1880, [m.], 3, 246. of. Cleve, Compt. rend., 1879, 89, 708
 Rescue, Trans. Chem. See, 1882, 41, 277; Crookes, Phyl. Trans., 1892, 174, 910.
 Urbain, Ann. Chim. Phys., 1906, [vii.], 19, 184.
 Delafontame, Chem. News, 1897, 75, 229
 Pelafontame, Compt. rend., 1878, 87, 632; 1880, 90, 221, Arch. Sci. phys. nat., 1880, [m.] 3, 250 Petatontaine, Compt. rena., 1819, 61, 322, 89, 212, of Delafontaine, loc. ett., and Compt. rend., 1878, 87, 634, Noret, ibid., 1879, 88, 422

10 Delafontaine, Compt. rend., 1881, 93, 63.

The discovery of samarium was confirmed by a number of chemists, and in 1883 Cleve prepared numerous samarium salts and determined the atomic weight of samarinin.1

In 1885 Aner von Welsbach made the remarkable discovery that "didyminm," even when freed from "samarinin," was complex, being a mixture of two elements which he named praseodymium and neodymium 2 The salts of these two elements were found to differ strikingly in colour, being green and rose-red respectively

In 1892 Lecoq de Borsbaudian discovered that Cleve's "samaria" was complex, and that when fractionated with ammonia, the least basic fractions differed from the others in two ways. (i) their spark spectra contained three new lines in the blue, \$4593 being the most conspicuous, and (ii) a new band \$46110-6220 appeared in their "spectres de renversement". The new spack lines he attributed to an element which be provisionally called  $Z_n$  and the band he similarly attributed to an element  $Z_n$  expressing no opinion as to the identity or otherwise of  $Z_n$  and  $Z_n$ . In 1893 he showed that  $Z_n$  was closely connected with an element  $S_n$ , the existence of which had been previously assumed by Crookes to account for the "anomalous band" encountered in the cathodic phosphorescence spectra of the rare earlies 4

The presence of a small amount of a new earth in "samaria" was further proved by Demarçay in 1896. Demarçay characterised his new earth by its ultraviolet spark spectrum, which was very sensitive, and provisionally called the new rare earth element Z In 1900 he showed that Z, Z, and Z were probably identical, a conclusion be confirmed in 1901, when he isolated the oxide of  $\Sigma$  in a nearly pure state. He then established the fact that  $\Sigma$  is identical with Crookes' S , and called the element  ${\bf europium}^{(n)}$ 

(m) The Terbeum and Mirrum Groups In 1878 Mangua Inschonated the seedolimite earths,7 and Soret examined the "erbia" mactions spectroscopically. As the result of his observations, Soret concluded that "erbin" was complex. Reserving the name of crbium for that rare earth element the salts of which gave rise to the most characteristic absorption bands of the old erhunn spectrum, Soret designated by X a new element, salts of which exhibit a characteristic absorption spectrum of some ten or more bands, particularly  $\lambda 6404$  and  $\lambda 5363$  . He also observed that a bind,  $\lambda 6840$ , which did not form part of the spectrum of N, did not appear to belong to the crbium spectrum either

Pursuing the fractionation of "crbia" by Berlin's method, Marignae in 1878 made the surprising discovery that the least basic portion of "erbia" consisted of a new, colomiless earth of very high chemical equivalent. This new earth he called oxide of ytterbium, " its spark spectrum was mapped by Lecoq de Boisbaudran 10

<sup>1</sup> Cleve, Compt. rend., 1883, 97, 94.
2 And von Welshach, Monatch., 1-95, 6, 177, et. that. 1884, 5, 1-508. As a matter of fact, von Welshach proposed the names praseodidymium and neodidymium, but the shorter rames given in the text are generally employed.

shotte terms given in the text are generally employed

Level de Borshaudran, Compt. rend., 1892, 114, 575, 1893, 116, 611, 674

The work of Crooker is discuss a later (p. 294)

Denortys, Compt. rend., 1896, 122, 728, of clot., 1886, 102, 1551, 1393, 117, 163.

Denortys, thet., 1900, 130, 1169, 1901, 132, 1484.

Minguac. Arch. Sci. phys. nat., 1878, 61, 283, Jan. Chim. Phys., 1878, [v.], 14, 247.

Soirel, Arch. Sci. phys. nat., 1878, 61, 283, Jan. Chim. Phys., 1878, [v.], 14, 247.

Manguac. Arch. Sci. phys. nat., 1878, 64, 97, Compt. rend., 1878, 86, 1662

Manguac. Arch. Sci. phys. nat., 1878, 64, 97, Compt. rend., 1878, 87, 578, acc. also
Delafontim. Compt. rend., 1878, 87, 933

Lecoq de Borsbaudran, Compt. rend., 1879, 88, 1342

NOL. 1897.

In the following year Nilson, who had prepared large quantities of "erbia," repeated and confirmed Marignae's work, but, in addition, he isolated still another earth from "crbia." It was colourless and less basic than ytterbia, and received the name of scandia or oxide of scandium. The same year Cleve I fractionated "erbia" freed from scandia and "ytterbia," using Berlin's method, and from the results of Thalen's spectroscopic study of the fractions, concluded that "crbia" was composed of three earths, which he called the oxides of holmium, erbium, and thulium in descendmg order of basicity Each earth was characterised by the absorption spectrum of its salts, and Soret and Clove quickly noted that holimum and Soret's X were identical, and that thulium was responsible for the somewhat anomalous band \$6840 in the spectrum of old "erbium"

The elementary nature of "holmum" was disproved in 1886 by Lecoq de Borsbaudran, who showed that its absorption spectrum characterised not one, but two elements Reserving the name of holimum for the element giving use to the characteristic bands \$\lambda 6104\$ and \$\lambda 5363\$ of Soret's X, he called the other element, which was defined by the absorption bands  $\lambda 7530$ and \$1515, dysprosium.5

It has been already mentioned (p. 223) that Mangnae confirmed the presence of "terbia" in the yttrium earths from gadolimte "Terbia" is more abundant in samarskite than in gadolinite, and in 1880 Marignac fractionated the samarskite earths by means of the double potassium sulphates He found 6 that "terbia" in samaiskite is accompanied by two new earths, which in properties approached the cerito rather than the gadolimite earths The two new elements he provisionally named  $\mathbf{Y}_{a}$  and  $\mathbf{Y}_{b}$ . It was considered highly probable by Marignae himself that Ya was identical with Lecoq de Boisbandran's samarium, discovered in the preceding year, and this opinion was confirmed by Soret  $^7$ . The fact that  $Y_\rho$  was a new element was confurred. · by Lecoq de Borsbaudran, and in 1886 Marignac named it gadolinium. Subsequently, Locoq de Boisbaudran found that Marignac's "gadohma" contained about 10 per cent of impurities, purified the material, and determined the atomic weight and spark spectrum of gadolimium

By the year 1886, then, Mosander's yttrum, erbinin, and terbinin had given place to the following list of elements Yttiium, gadoliuum, terbium, dysprosium, holmium, erbinin, thininin, ytterbinin, and scandinin. In addition to these elements, three other provisional elements,  $Z_a$ ,  $Z_b$ , and  $Z_n$ , had been announced by Lecoq do Borsbaudian as being present in "terba," in and the nature of "mosandrum," "deepnum," and "phillipium" was still in doubt. Moreover, in 1886, Crookes propounded his celebrated theory of meta-elements as an interpretation of the results of several years

Nilson, Compt. rend., 1879, 88, 645, Ber., 1879, 12, 551

See Chapter IX

Cleve, Compt. rend., 1879, 89, 478, Chem. News, 1879, 40, 125,

Soict, Compt. rend., 1879, 89, 521, Cleve, ibid., 1879, 89, 708, Lecoq de Boisbaudran, ibid., 1878, 89, 516

Lecor de Borsbaudian, Compt lend, 1886, 102, 1003, 1005.

Mangnac, Arch. Sci. phys. nat., 1880, [m], 3, 413, (ompt. rend., 1880, 90, 899;
 Ann. Chm. Phys., 1880, [v], 20, 535
 Smet. Compt. rend., 1880, 91, 378, Arch. Sci. phys. nat., 1880, [m], 4, 261.
 Announced by Locoq de Boisbaudian to the French Academy, Compt. rend., 1886, 102, 00.

<sup>02.</sup>Lecoq de Borsbaudran, *ibid* , 1889, 108, 165 ; 1890, III, 393, 409, 472.

<sup>10</sup> See p. 227.

of research on the cathodic phosphorescence spectra of the rare earths.\(^1\) At that time, therefore, the chemistry of the rare earth elements appeared to be of the most amazing complexity, and the confusion was increased when in 1887 Kruss and Nilson put forward a "one band -one element" theory of the absorption spectra of the rare earths \(^2\)

At the present time the chomistry of the rare earths is in a much more satisfactory state. The theories of Crookes and of Kriss and Nilson have been abandoned, and quite a number of rare earths which in 1886 were only recognised as distinct bodies from spectroscopic observations have now been isolated in a state of purity. Thus, pure gadolima and terbia were isolated in 1905 by Urbain, and the isolation of pure dyspressa was effected by the same chemist in 1906. Practically pure holima was isolated in 1911 by Holinberg, and thulia by James in the same year. Pure erbia, however, has not yet been prepared, though nearly pure preparations have been obtained by Urbain, James, and K. A. Hofmann, lastly, "ytterbia," has proved to be complex.

The complexity of "ytterbar" was discovered independently by Aner von Welsbach and Urbain. The former chainst amounted his discovery in 1905, and termed the component elements of "ytterbium" aldebaranium and cassiopeium. The latter, who in 1907 published preliminary measurements of atomic weights and spectra, named the component elements neoytterbium and lutecium. The International Committee on Atomic Weights has adopted the names ytterbium and lutecium respectively.

In 1911 Urbain's amounted that the luterium obtained from gudolimite is accompanied by a small amount of still another rare earth element, which he named celtium

The previously mentioned elements  $Z_{\alpha}$   $Z_{\beta}$ , and  $Z_{\gamma}$  were discovered spectroscopically by Lecoq de Boisbaudian  $Z_{\alpha}$  was characterised by the bands  $\lambda5730$  and  $\lambda4765$ , and  $Z_{\beta}$  by the bands  $\lambda6205$ ,  $\lambda5858$ ,  $\lambda5132$ , and  $\lambda4870$  in its "spectre de renversement"."  $Z_{\gamma}$  was defined by the lines  $\lambda5835$ ,  $\lambda5750$ ,  $\lambda5700$ ,  $\lambda5269$ , and  $\lambda5259$  in its spark spectrum in In 1886 Lecoq de Boisbaudian also concluded that a fourth unknown element,  $Z_{\delta}$ , was present in "terbin" and characterised it by the absorption band  $\lambda4880$ , but this deduction was not published until 1895 ii

In 1900 Demarcay 12 announced four more "spectroscopic" elements,

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    See later, p. 295.
    See later, p. 287.
    Itham, Compt. rend., 1905, 140, 583 (Gd., 1995, 141-521 (Th)
    Urbam, Compt. rend., 1905, 140, 583 (Gd., 1995, 141-521 (Th)
    Urbam, Lind., 1906, 142, 785
    Holmberg, Artiv Kem. Min. Good., 1911, 4, Nos. 2 and 10
    Lames, J. Amer. Chem. Soc., 1911, 33-1332
    Auer von Welsbach, Anzener K. Akad. Wiss. Wien, 1905, No. 10. Innaten, 1907, 351, 464, Monatch., 1906, 27-95, 195, 1908, 29, 181, Scieunsber K. Akad. Wiss. Wien, 1906, 175, II. B, 737, 1907, 116, II. B, 1425; Urbam, Compt. rend., 1907, 145, 759.
    See also. Urbam, Chem. Zeit., 1908, 32, 730, Zeitsch. amorg. Chem., 1910, 68, 232, Auer on Welsbach, Sitzungber. K. Akad. Wiss. Wien, 1909, 118, II. B, 507, Monatsh, 1909, 30, 695.
    Wenzel, Zeitsch. amorg. Chem., 1909, 64, 119
    Urbam, Compt. rend., 1911, 152, 141.
    Levon de Boisbaudran, Compt. rend., 1885, 100, 4437, 101, 552, 558, 1889, 102, 395, 483, 899, 1536, 103, 113, 627, 1887, 105, 258, 301, 344, 784, 1890, 110, 21, 67, 111, 102 Levon de Boisbaudran, Compt. rend., 1885, 102, 153.
    Levon de Boisbaudran, Compt. rend., 1885, 102, 159.
    Demarçay, Compt. rend., 1900, 131, 387.
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 $\Gamma$ ,  $\Delta$ ,  $\Omega$ , and  $\Theta$ , each being characterised by its ultraviolet spark spectrum. According to Demarçay,  $\Gamma$  was possibly identical with true terbium,  $\Delta$  was probably  $Z_{\gamma_1}$   $\Omega$  (defined by the lines 3967 9 and 3930 9) came between holimon and erbium in his fractions, and  $\Theta$  (defined by the lines 4008 2 and 2006 5) between and attrabases. 3906 5) between erbium and ytterbium.

Of the eight elements,  $Z_a$ ,  $Z_b$ ,  $Z_\gamma$ ,  $Z_b$ ,  $\Gamma$ ,  $\Delta$ ,  $\Omega$ , and  $\Theta$ , only two, namely  $\Omega$  and  $\Theta$ , still await isolation or identification with other elements, for Urbain has shown conclusively that  $Z_{\beta}$ ,  $Z_{\delta}$ , and  $\Gamma$  are identical with terbium, and that  $Z_{\alpha}$ ,  $Z_{\gamma}$ , and  $\Delta$  are identical with dysprosium  $^{2}$  (iv) Summary.—The history of the discovery of the elements of the rate earths is summarised in the accompanying table.— $^{3}$ 

| Date.        | Element         | Material in<br>which it was<br>discovered | Discoverer                    | Remarks                      |
|--------------|-----------------|-------------------------------------------|-------------------------------|------------------------------|
| 1791<br>1804 | Yttmum<br>Cemum | Gadolinite<br>Cerite                      | Gadolin<br>Berzelius and Hism |                              |
| 1839         | Lanthanum       | Cern                                      | ger , Klaproth<br>Mosander    | 1                            |
| 1842         | Didynnum        | 1                                         | 1                             |                              |
| 1843         | Terbium .       | Yitma                                     | "                             | 1 (1-11-1                    |
|              |                 |                                           | **                            | Called cibium from           |
| 1843         | Erbium .        | 1 ,,                                      | 1                             | 1860 onwards                 |
|              |                 | . ,,                                      | <b>,</b> ,,                   | Called terbium from          |
| 1878         | Mosandrum       | Samuskite                                 | Liwience Smith                | 1877 onwards                 |
| 1878         | Phillipum       | Yttina carths                             | Delatontaine                  | A mixture                    |
| 1878         | Decipium .      | Samuskite                                 | Delatoritaring                | . ,                          |
| 1878         | Ytterbium       | Etbia                                     | Manguac                       | ,,,                          |
| 1878         | λ.              | 1                                         | Solet                         |                              |
| 1879         | Scandrum        | Ytterbia                                  | Nilson                        | Cleve's holmman              |
| 1879         | Samarum         | Didyima                                   | Lecoq de Boisbaudian          | • 17                         |
| 1879         | Thulum          | E bia                                     | Cleve                         |                              |
|              | Holmun          |                                           | (10.0                         | Isolated in 1911             |
| 1880         | Ya .            | Samarskite                                | Marignac                      | Named gadolinium ii          |
| 1885         | M I             | 1                                         |                               | 1886                         |
| 1885         | Neodymnum       | ' Didynna                                 | Auer von Welsbach             | 1                            |
| 1885         | Prascodymium.   | i m . 11                                  | ,,                            |                              |
| 1885         | Za .            | Terbia carths                             | Lecoq de Borsbaudran          | Dysprosium                   |
| 1886         | Ζβ .            |                                           | • ,,                          | Terbrum                      |
| 1886         | Zγ              | ' "                                       | ,,                            | Dysprosium                   |
|              | Z <sub>δ</sub>  |                                           | ;,                            | Terbmin                      |
| 1886         | Dysprosium      | Holma                                     | .,                            | Isolated in 1906             |
| 1886         | Meta-clements   |                                           | Crookes                       | See p 295                    |
| 1887         | ., ,,           | ~                                         | Kruss and Nilson              | Sec p. 287                   |
| 1892<br>1892 | Z               | Samaria                                   | Lecoq de Borsbandran          | Europium                     |
| 1892         | Ζζ .            | ,,,,                                      |                               | 11                           |
|              |                 | Yttiia carths                             | Rowland                       | Dy sprosium See p            |
| 1896         | <b>x</b>        | "                                         | Demarçay                      | Europium Isolated<br>in 1900 |
| 1896         | Lucinm .        | Yttua from                                | Barrière                      | Amixture Seep 363            |
| 1896         | Kosmium .       |                                           | Kosmann                       | ,, ,,                        |

<sup>&</sup>lt;sup>1</sup> Urban, Compt. rend., 1905, 141, 521, J. Chem. phys., 1906, 4, 334; Lecoq de Bonsbaudran, Compt. rend., 1904, 139, 1015

<sup>2</sup> Urban, Compt. rend., 1906, 142, 785, J. Che. phys., 1906, 4, 356.

<sup>3</sup> See Baskerville, Chem. News, 1904, 89, 150.

| Date.  | Element              | Material in<br>which it was<br>discovered | Discoverer         | Remarks.                            |
|--------|----------------------|-------------------------------------------|--------------------|-------------------------------------|
|        |                      |                                           |                    | -                                   |
| 1896   | Neokosimum           |                                           | Kosmann            | A mixture. Seep 36                  |
| 1897   | Glaukodyminin        | Didymia                                   | Chroustschaft!     | Nature unknewn                      |
| 1899   | Victorium<br>Omonimo | Ytti a cuths                              | Crookes            | Disployed by Urbain.<br>Scr. p. 301 |
| 1900   | Г                    | . Toltreanths                             | Penarcay           | Terbum                              |
|        | Δ                    | •                                         | •                  | Dysprosium                          |
|        | Ω                    | ( Yithiy carths                           |                    | Unknown                             |
|        | H                    |                                           |                    |                                     |
| 1907   | Incognitum           |                                           | Crookes            | Perbum See p 30:                    |
| 1905   | Immuno               | •                                         |                    | P 200                               |
| 1905   | Neoviterbum          | Yukuban                                   | Anci ven Welsbach, | ու շուր ասա                         |
| 1907   | (aldebaranenn        |                                           | Urbain             |                                     |
| 1405   | Lutecium             |                                           | 1 1141111          |                                     |
| 1907   | (*85510]** 111111    | **                                        | •                  |                                     |
|        |                      | Lutina                                    | Urbam              | 1                                   |
| 1911 , | Celtiniii            | Luteri                                    | CIDOIN             |                                     |

# THE RARE EARTH METALS

Preparation Few of the rare cartb metals have been isolated in a state or purity. The following methods have been used -

(1) Reduction of the anhydrous chloride by heating it with sodium or potassium. Most experimenters have found it very difficult to obtain even a moderate yield of metal by this process. The metal is very hable to be produced in a large number of time globules, difficult to remove from the alkali chlorale produced further, the metal is hable to contain sodium or potassium

(ii) Reduction of the authydrous fluoride by heating it with calcium or aliminium. This method does not appear to have firmshed the pure metals, but cerium calcium and cerium aluminium alloys have been thus obtained.8

(iii) Reduction of the oride by heating it with magnesium, almemium, 5 calcium, carbon, or silicon. Pure rare carth metals have not been obtained by any of these means. When magnesium, aluminium, or calcium is employed, alloys are usually obtained while with aluminium there is evidence of the production of subovides of the rare earth elements. When reductions

<sup>&</sup>lt;sup>1</sup> Chroustschoff, J. Russ. Phys. Chem. Soc., 1897, 29, 206.

<sup>2</sup> Mosander, Poor. Annalen, 1826, 11, 109, 1839, 46, 648, 47, 207, 1843, 56, 504; Annalen, 1849, 32, 235, 1843, 48, 210, Phil. Wats, 1843, 23, 241 (Ce., Let., Ch.), Bernger, Annalen, 1842, 42, 134 (Ce), Berreines, Lehrbach (1826, vol. 2), p. 416, Wohler, Poop. Annalen, 1828, 13, 580 (Y), Annalen, 1867, 144, 251 (Ce), Poop. Landen, 1864, 131, 179 (Y), Margass, Ann. Chim. Phys., 1853, [n], 38, 148 (Ph.), Gove and Hogland, Bull Soc. chim., 1873, [n], 18, 189 (Y), Cleve., ibid., 1874, [n], 21, 341 (Y); Matignon, Ann. Chim. Phys., 1906, [vin.], 8, 282 (Nd); Hunter, Eighth Inter. Cong. Appl. Chem., 1912, 2, 125 (Nd)

<sup>1</sup> Moble inhauer, Chem. Zeid., 1914, 38, 147

<sup>4</sup> Winklet, Bei., 1820, 23, 772, 1821, 24, 573 (La, Y, Ce), Matignon, Compt. vend., 1960, 131, 837.

<sup>8</sup> Matignon, loc. cit., Schiffer, Inaugural Inscription (Munchen, Tech. Hechschole).

<sup>1900, 131, 831.</sup>Mangnon, loc cit., Schiffer, Inaugural Dissertation (Munclen, Tech. Hechschule);
Moldenhauer, loc cit., Hirsch, J. Ind. Eng. Chem. 1911, 3, 880, 1912, 4, 65, Trans.
Amer. Flectrochem. Soc., 1911, 20, 57.

Hirsch, loc. cit.

with carbon or silicon are carried out, the products are carbides and silicides of the rare earth elements

- (iv.) Electrolysis of the molten, anhydrous chlorides, with or without the addition of the chlorides of sodium, potassium, calcium, or barrum 1. The chloride may be melted in an iron crucible and electrolysed, using carbon electrodes In carrying out the process, by means of which considerable quantities of the rare earth metals of the cerum group have been obtained, numerous practical difficulties arise, so that it is not possible usefully to fore referred to the original memoirs of Hirsch and of Muthmann and his co workers 2
- (v) Electrolysis of the molten fluoride, or better, a solution of the oxide in the fluoride 3. According to Muthmann and Scheidmandel, this process is superior to the previous method, but Hirsch holds the opposite view

The rare earth metals obtained by electrolysis may be purified by melting them under a layer of barum chloride in a magnesia crucible

Properties 1 -The rare earth elements are lustrous metals has the whiteness of tin, cerium that of iron, needymium has a yellow tinge, and praseodymum is decidedly yellow, samarium is pale grey in colour With the exception of lanthamim, they retain their lustre in dry air. When compared with lead, tin, and vine, the order of mereasing hardness is as follows Pb, Su, Ce, La, Zu, Nd, Pi, Sm. Commit can easily be cut with a knife, samarium is as hard as steel. The densities and melting-points are given by Muthmann and Weiss as follows -

|                   | L.,      | Ce   | Pı    | Nd.   | Sm.     |
|-------------------|----------|------|-------|-------|---------|
| Melting-point, *C | 8107     | 623° | 940°  | 840°  |         |
| Density at 20° C  | $6\ 155$ | 7012 | 6.475 | 6.956 | 7 727 8 |

Husch gives the values 635° and 6 92 (at 25°) for corum.

The nare earth metals burn in air or oxygen very readily, with the evolution of much heat and light (p 257) Cerum, for example, begins to burn at 160° m air. They also combine with hydrogen, nitrogen, chlorme, and bromme when heated in those gases, and react with iodine, sulphur, phosphorus, etc. They are, in fact, highly reactive elements. In dilute mineral acids they readily dissolve, and, except with nitite acid, hydrogen is evolved They are also attacked by water, very slowly at ordinary temperatures, but more rapidly at 100

Numerous cerrum alloys and a few alloys of the other rare earth metals have been prepared: one of these may be mentioned here, viz. the alloy known as mischmetall. Owing to the large amount of heat evolved when any of the rare earth metals is burnt in air, these metals form excellent

<sup>&</sup>lt;sup>1</sup> Erk, Zeitsch fur Chem., 1870, [n.], 7, 100, Jahresber, 1870, p. 319 (Ce); Cleve and Hoglund, Bull Suc chim, 1873, [n.], 18, 193, 289 (Y); Frey, Annalen, 1874, 183, 367 (Ce); and particularly the following Hillebrand and Noton, Popp Annalen, 1875, 155, 633; 156, 466 (Ce, La, "Di"), Muthmann, Hotel, and Weiss, Annalen, 1902, 320, 231 (Ce, Md); Mu'hmann and Kraft, Annalen, 1902, 325, 261 (Ce, La), Muthmann and Weiss, ibid., 1601, 331, 1 (Ce, La, Nd, Pi, Sm), Hirsch, loc. cit. (Ce), Borchers and Stockem, D.R.P., No. 172,529.

§ Kull kitzle, Mulligher, Will who be found in Kellanuann, Die Contractale und also more more contractal.

D.R.P., No. 172,529.

8 Full details will also be found in Kellermann, Die Ceritmetalle und ihre pyrophoren Legierungen (Knapp, Halle, 1912)

Muthmann and Scheidmandel, Annalen, 1907, 355, 116; Hirsch, loc. ett.

Hillebrand and Norton, loc. est., Muthmann and others, loc. ett.; Hirsch, loc. ett.

reducing agents for preparing other elements from their oxides. It is abvious that a mixture of rais earth metals would do equally well for this purpose; hence the preparation of mischinetall, which consists essentially of cerium, lanthanum, prascodymum, and neodymum, and is prepared from the residual salts of these metals left over after the thorium and sufficient cerum and "didymnim" for the gas mantle industry have been extracted from morazite sand Mischnetall reduces the oxides of mimerous elements, eg non, cobalt, nickel, iornganese, chromium, molybdenim, vanadium, columbuno, tantahun, silicon, boron, tin, lead, titannim, and zirconnim, and has been of considerable service in the preparation of a number of rare olements 1

# VALENCY, ATOMIC WEIGHTS, AND CLASSIFICATION OF THE RARE EARTH ELEMENTS.

Theoretical.<sup>2</sup> Historical Resume -In 1810 the rare earths had only been separated into "ceria" and "vttria," and Berzehus, in constructing his first atomic weight table, regarded yttim as YO<sub>9</sub> cerm as CeO<sub>9</sub> and the lower oxide (cerons oxide) as CeO — In 1826 he first admitted the exist nee of sesquioxides, and in the changes nocessitated in his atomic weights and formula the three preceding formula were involved and became YO, Ce,On and CeO respectively.

By the year 1842 the rare earths had been separated into cerra, lanthana, "didymia," yttiia, "crbin, and "terbia", except for the fact that Mosander's "erbia" was denied existence by various chemists, their number remained imaltered until 1878, and prior to 1870 then formula were all written in the form MO, ceria being excepted. These formula were chosen because the raic Aiths were strong bases like line, strontia, and baryta, and on Gmelin's system of atomic weights (the one most commonly adopted) these substances were written CaO, SiO, BaO Ceria, however, was discovered to be Ce<sub>a</sub>O, if cerous oxide is CeO, and not Ce<sub>2</sub>O<sub>4</sub> as had previously been supposed , and for some years it was supposed that ceria was a compound of two oxides CeO and CeO, (Ce O, being 2CcO CeO,), the oxide CeO, being the one from which ceric salts were derived. The latter assumption was not difficult to make, for pure ceric salts had not been prepared and analysed

The evolution of the modern system of atomic weights in 1858 did not affect the formulae of the compounds of the rare earth elements, for the necessary experimental data were not then available. The proposal to change the accepted atomic weights and formula was made in 1870 by Mendeldeff, who considered that the following formulae should replace those that were then in use -

Valuable support was soon forthcoming for Mendelécil's views In 1872 Cleve agreed that the tervalency of yttrmm and erbum was very probable, and pointed out the fact that the formula of their compounds were thus

See Wei-s and Aichel, Annale t, 1994, 337, 379
 Throughout this section, cere compounds are disregarded, except where the contrary is expressly stated

<sup>3</sup> Mendeléeff, Annalen Suppl., 1872, 8, 133, translation in Chem. News, 1879, 40, 231, etc.; 1880, 41, 2, etc., see also Mendeléeff, Bull. Acad. Sci. Petrograd, 1870, p. 415.

simplified. Two years later, he definitely accepted the tervalency of yttrium, erbium, "didymium," and cerium (in the cerous salts), but differed from Mondeléeff in regarding Luthanum also as a triad 1 In 1876 Hillebrand showed that the specific heats of lanthanum, cerium, and "didymium" were in harmony with Cleve's view,2 and Rammelsberg,3 who had previously objected to Mendel/eff's proposals,4 accepted those of Cleve.

From that time onward the tervalency of the rare earth elements has been generally accepted, occasionally, however, a dissentient voice has been heard Thus, Cossa 'argued for the bivalency, and Winkler for the quadrivalency of these elements. Winkler, however, abandoned his views after they had been criticised by Brauner, 7 and it has iemained for only two chemists—Wyronboff, and his frequent collaborator, Verneuil—consistently to uphold the bivalency of the rare earth elements 8. These two French chemists decline to accept the physical evidence of specific heats and isomorphism, the latter on famly good grounds, and utterly reject any considerations based upon the Periodic Classification, which they regard as a metaphysical conception. Instead, they demand chemical proofs, and consider from that point of view that they can establish a very close relationship between the rare each elements and the bivalent metals. Into the nature of these chemical relationships, however, it is hardly necessary to enter, for, whatever value may attach to them, Urbain has shown that similar relationships may be recalled which connect the rare earth elements at least as well with bismuth 10

The evidence bearing on the valency of the rare earth elements will now be considered in some detail

(1) Dulong and Petit's Law - The only specific heat data for the elements themselves are given in the following table -

| Element     | Specific Heat         | Temp<br>Range       | Combining<br>Weight | Combining Weight's<br>Specific Heat |
|-------------|-----------------------|---------------------|---------------------|-------------------------------------|
| Cermm       | 0 0448 11<br>0 050 12 | 0, 100,             | 46.8                | 2 09 -6 4"3 approx<br>2 31 -6 4/3   |
| Lanthanum . | 0 0511 13             | 20' 100°<br>0' 100° | 46 8<br>16 3        | 2 39-6 4/3                          |
| "Di tymmm"  | . 0 0456 11           | 0°- 100°            | 17 7                | 2 18 6 1/3 ,,                       |

Cleve and Hoglund, Bull Soc chim., 1872, [n], 18, 193, 289; Cleve, ibid., 1874, [n], 21, 196, 216, 344.
 John, ibid., 1874, [n], 21, 553.
 Hillebrand, Topp. Innaten, 1876, 158, 71.
 Rinnielsberg, En., 1876, 9, 1580.
 Rammelsberg, En., 1873, 6, 84.
 See also Delafontame, Arch Sci. phys. nat., 1874, 58ce, 1234.
 See p. 234.

<sup>51, 45,</sup> and Mendeléeft, Annalen, 1873, 168, 45.

<sup>2</sup> See p. 234

<sup>6</sup> Winkher, Ber., 1891, 24, 873

<sup>7</sup> Braunet, Ber., 1891, 24, 1328, Winkley, Ber., 1891, 24, 1966

<sup>8</sup> See particularly Wyrouboff, Bull. Soc. Irang. Min., 1896, 19, 219, 1905, 28, 201, Wyrouboff and Verneuil, Bull. Soc. chim., 1899, [iii], 21, 118, Ann. Chim. I hys., 1905, [viii], 6, 465-482.

<sup>9</sup> See p. 236.

<sup>10</sup> Urbain, J. Chim., phys., 1906, 4, 107.

<sup>11</sup> Hillehrand, Pogg. Annalen. 1876, 158, 71.

<sup>12</sup> Mendeléeft, Bull. Acad. Sci. Pebograd, 1870, p. 445.

<sup>13</sup> Hirsch, J. Ind. Eng. Chem., 1911, 3, 880, 1912, 4, 65.

It will be seen that the results argue very strongly for the tervalency of the elements

The extension of Dulong and Petit's Law to compounds, usually called Kopp's Law, may also be utilised. The mean specific heats between 0° and of a number of the rare earths and their sulphates were measured by Nilson and Pettersson, from whose memorr the following figures are

| Compound                                                                                                  | Specific<br>Heat ,                                                                 | Molecular<br>Heat                                                             | Componed                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | Specific<br>Heat                                                                                  | Molecular<br>Heat                                                                                        |
|-----------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------|-------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------|
| Al 0<br>Se 0<br>Ga,0,<br>V <sub>2</sub> 0.<br>In 0<br>Et 0,<br>V <sub>1</sub> 0,0,"<br>L <sub>1</sub> 0," | n 1870<br>n 1840<br>n 1062<br>n 10 of<br>n 0 0807<br>n 0 0816<br>n 0749<br>n 0 810 | 19 32<br>29 81<br>19 51<br>23 29<br>22 17<br>24 70<br>25 45<br>21 42<br>27 62 | M <sub>1</sub> (SO <sub>4</sub> )   1<br>Se <sub>4</sub> (SO <sub>4</sub> )   1<br>Ge <sub>4</sub> (SO <sub>4</sub> )   1<br>Y (SO <sub>4</sub> )   1<br>H <sub>1</sub> (SO <sub>4</sub> )   1<br>H <sub>2</sub> (SO <sub>4</sub> )   1<br>H <sub>3</sub> (SO <sub>4</sub> )   1<br>Ge <sub>4</sub> (SO <sub>4</sub> ) | 0 (855)<br>0 16,9<br>0 1460<br>0 1799<br>0 1290<br>0 1069<br>0 1182<br>0 1187<br>0 1168<br>0 1656 | 61 59<br>62 47<br>61 90<br>61 bu<br>66 41<br>61 48<br>65 87<br>66 10<br>68 96<br>68 24<br>66 24<br>67 41 |

The tervalency of the rare earth elements mentioned in the preceding table is thus seen to be in harmony with Kopics Law, when the molecular heats of the oxides and sulphates of the tervalent elements aliminium, scandigm, gallium, indium, iron, and chromium are compared with the similar data for the compounds of the rare earth clements. The evidence, however, is not very strong. From measurements of the specific heats of cerous and "didymium" tungstates Cossa concluded that the rare earth elements are diads 4

(n ) Transpurency to X-rays Benoist's method, as described under Indium (see pp. 152-1), has been applied to cerum, and the results obtained show that the atomic weight of the element is three times the chemical equivalent of cerum in cerus salts?

(m) X-ray Spectra —The X-ray Spectra of the elements are discussed later in this volume (see pp. 312, 363), where it is shown that the atomic numbers of the rare earth elements (yttrium excepted) he between those of barmin and tantaliin. Since, in general, the order of increasing atomic numbers is the same as that of mereasing atomic weights, it follows that the atomic weights of the rare earth elements he between those of barrum and tantalum, a clear indication that the rare earth elements are tervalent

(iv) The Law of Isomorphism - It very frequently happens that corresponding compounds of the rare earth elements are isomorphous. The actual cases need not, however, be cited here, as full details are given later

Nilson and Pettersson, Compt. rend., 1880, 91, 232; Rev., 1880, 13, 1459, 27. Tanatar and Volpusky, J. Ruse. Thus. Chem. Soc., 1910, 42, 39.

It may be remarked that the atomic heats of the chaments aluminum and gallium are decidedly below the normal value 6.4. The values for scandium and yttrium are not known.

Costa and Consus. Costa and Zichini, ride inforce.

Remove and Consus. Costa and Zichini, ride inforce.

<sup>&</sup>lt;sup>8</sup> Benoist and Copaux, Compt. rend , 1914, 158, 689

in this chapter, when the rare earth salts are described in some detail

(see pp 250-282).

The conclusion that may be drawn from the preceding fact is very important. The ratio atomic weight : combining weight is the same for all the rare earth elements, the combining weights being deduced from the compositions of the basic oxides. The determination of this ratio by utilising the law of isomorphism cannot, however, be effected unless cases of isomorphism between compounds of the rare earth elements and of other elements of known atomic weight can be discovered. A number of such cases are actually known, and the conclusions to which they lead are conflicting; in part the results indicate that the rare earth elements are diads, and in part they point to their being triads

(a) Cerous and "didymium" tungstates and molybdates are isomorphous, and form mixed crystals with calcium tungstate, CaWO4, and lead molybdate, PbMoO.

(b) Calcium and strontium silicoting states are isomorphous and form mixed crystals with the subcotning states of lanthanium, ceruim, and "didy mium"?

(c) Lanthanium, ceruin, and "didymium" silicotning states are also iso-

morphous with the thorum salt 3

(d) In certain numerals, e.g. parisite, cordylite, yttrocerite, and yttrofluorite, the fluorides of calcium, barrum, and the rare earth elements are present in isomorphous mixture 1

(e) Bisimuth intrate forms a stable pentahydrate,  $Bi(NO_3)_a$   $5H_2O_5$  and a labile hexahydrate, Bi(NO3) 611,0, lanthanum and "didymium" intrates form stable herahydrates, M(NO<sub>3</sub>), 611,0, and labile pentahydrates, M(NO<sub>3</sub>), 511,0, if the metals are regarded as tervalent, and these salts are isodimorphous with bismuth nitrate, two series of mixed crystals being formed with the La-Bi and two with the "Di"-Bi mixtures Two series of mixed crystals are also formed with the yttrium bismuth mixtures?

Bismuth intracte,  $B_1(NO_3)_q$   $5H_2O_1$  is isomorphous with the intractes of the metals of the terbiam and yttrium groups obtained by crystallising from fairly concentrated intrie acid (density 1.3). These hydrates are of the type MIII(NO<sub>3</sub>), 511,0 if the rare earths are tervalent. The gadolimum, terbium, dysprosium, and holimum salts form mixed crystals with the bismuth salt, which also induces the crystallisation of neoytterbium and lutecium initiates

The hydrates of bismuth sulphate are isomorphous with various hydrates of the rare earth sulphates, and the following mixed crystals have been prepared, but only containing a few per cent of the rare earth salts --

$$\begin{array}{ll} (B_{1},La)_{2}(SO_{4})_{3} \; 9H_{2}O \\ (B_{1},La)_{2}(SO_{4})_{3} \; 5H_{2}O \end{array} \qquad \begin{array}{ll} (B_{1},"\; D_{1}")_{*}(SO_{4})_{4} \; 8H_{2}O \\ (B_{1},Y)_{2}(SO_{4})_{3} \; 8H_{2}O \end{array}$$

Cossa, Gazzetta, 1879, 9, 118; 1880, 10, 167, Compt. rend., 1881, 98, 900. 1886. 102, 1315; Cossa and Zecchini, Unzzetta, 1880, 10, 225, Dulier, Compt. rend., 1826, 102, 823; Hogbom, Bull. Soc. chem., 1884, [11], 42, 2
 Wyroubodl, Bull. Soc. franç. Min., 1896, 19, 219; 1905, 28, 201.
 Wyroubodl, Bull. Soc. franç. Min., 1896, 19, 219; 1905, 28, 201.
 Wyroubodl, Bull. Soc. franç. Min., 1896, 19, 219; 1905, 28, 201.
 Wyroubodl, Bull. Soc. franç. Min., 1896, 19, 219; 1905, 28, 201.
 Wyroubodl, Bull. Soc. franç. Min., 1896, 19, 219; 1905, 28, 201.
 Wyroubodl, Bull. Soc. franç. Min., 1900, 26, II., No. 3., Zeitsch. anong. Chem., 1901, 27, 254; Zeitsch. Kryst. Min., 1900, 32, 613, 1902, 36, 192, Ber., 1898, 31, 1227.

<sup>&</sup>lt;sup>8</sup> Urbain, Compt. rend. 1904, 139, 736, 1909, 149, 37, J. Chem. phys., 1906, 4, 342; Blumenfeld and Urbain, Compt. rend., 1914, 159, 323. 7 Bodman, loc. cit.

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Lanthanum ammonium nitrate, which is  $La(NO_2)_3 \cdot 2(NH_4)NO_3 \cdot tH_4O_5$  on the assumption that lanthanum is tervalent, forms mixed crystals with bismuth ammonium nitrate,  $\mathrm{Bi}(\mathrm{NO_2})_3$  2(NH<sub>4</sub>)NO<sub>3</sub> 4H<sub>2</sub>O.1

Bismuth nitrate forms double salts with the nitrates of magnesium, rinc, nickel, cobalt, and manganese. They are of the type  $2\mathrm{Bi}(\mathrm{NO}_s), 3\mathrm{M}^{\mathrm{H}}(\mathrm{NO}_s), 24\mathrm{H}_2\mathrm{O},$  hexagonal, and isomorphous with one another. The nitrates of lanthanum, cermin, praseodyminin, neodyminin, samarium, europinin, and gadolmum also form double intrates with the diad elements already mentioned. They are of the type  $2M^{11}(NO_s)_3 3M^{11}(NO_s)_2 24H_0O_s$  if the rare earth elements are triads. These salts are hexagonal and strictly isomorphous with the bismuth salts, the latter readily form mixed crystals with the salts of gadolmum, europeum, and samarium 2

Summing up the crystallographic evidence, it may be said that while the evidence given in puragraphs (a), (b), and (d) above is in favour of the view that the rare earth elements are diads, that given in (c) suggests that they are tetrids, and that given in (c) strongly suggests that they are triads. The evidence derived from the study of the silicotangstates cannot be considered at all conclusive. It may obviously be used to support the view that thorum is a daid, a view contrary to all the other evidence bearing on the valency of thorium, 3 and if may be added that it also leads to the untenable conclusion that lithium is a duid 4. In fact, with such a large molecule as that of a silicotung state, the nature of the metal present would appear to exert only a minor influence on the crystalline structure. The cases of the molybdates and tungstates, however, cannot be ruled aside in a similar manner, but in view of the strong evidence in favour of the tervalency of the rice earth elements afforded by then isomorphism with bismuth, it is possible that the cases of the molybdates and tungstates are to be ranked with the other "unusual" cases of isomorphism that have been pointed out by Barker? The safest plan, nevertheless, is to ignore the evidence based upon considerations of isomorphism

(v) Molecular Weight Measurements - No compounds of the rare earth clements are known that volatilise without decomposition at temperatures suitable for vapour density determinations. Molecular weights have accordingly to be measured in solution

(a) The anhydrous chlorides of lanthanum, cerum, prascodymum, neo dymnum, and yttrium have been studied in alcoholic solution by the horlingpoint method. The molecular weights are in accordance with the simple formulæ MulCl3, the metals being tervalent 6

(b) The chlorides of neodymnum, samarum, and "ytterbium" have been examined in aqueous solution by the freezing-point method. Here, owing to ionisation, it is necessary to express comparative results. For solu-

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Urbain and Lacombe cited by Urbain, J. Chim. phys., 1906, 4, 107
 Urbain and Lacombe, Compl. rend., 1903, 137, 508, 792, 820; Urbain, J. Chim. phys.,

<sup>1906, 4, 105.</sup>See Vol V of this series.

<sup>See Vol V of this series.
See Wyroubol, loc cil
Barker, Trans. Chim Soc., 1912, 101, 2484
Muthmann, Ber., 1898, 31, 1829 (Ce), Brauner, Proc. Chem. Soc., 1901, 17, 65;
Chem. Zentr., 1900, in 524 (Pr), Matignon, Ann. Chem. Phys. 1906, [viii], 8, 273 (Nd), 435 (Y), W. Biltz, Annalen, 1904, 331, 334 (La. "Dr"). In alcoholic solution the raise earth chlorides are undoubtedly combined with the solvent (see p. 255).</sup> 

tions which freeze at - 1° (' the following "molecular depressions" have been

| NdC1, |  | 68   | AICI,      |  | 64.5 |
|-------|--|------|------------|--|------|
| SmCI. |  | 65.6 | CiCl, .    |  | 65   |
| YbC1, |  | 66   | $Al(NO_i)$ |  | 65 4 |

The foregoing results are calculated on the assumption that the rare earth metals are tervalent, and the assumption is seen to be very satisfactory The alternative view that they are diads is much less satisfactory

(c) Various acetylacetonates of the rare earth metals have been examined in organic media. The results are in harmony with the view that the acetylacctonates have the double molecular weight, i.e. [Mai](CH, CO),(CH),[]] in moderately concentrated solution, and that the double molecules break down with dibition The following cases have been examined --

Ce compdom carbon tetrachloride, "Dr" compdom carbon tetrachloride. ,, ,, disulphide, ,, ,, ethyl sulphide. Sm " " " disulphide

(d) The equivalent conductivities of aqueous solutions of various salts of the rare earth elements have been measured. The following results have been obtained at 25° -- 1

According to Brauner and Švagr, the value of  $\lambda_{1021} - \lambda_{32}$  at 25° for each of the sulphates of lanthanum, cerum, praseodymum, neodymum, samarum, gadolinium, vttimm, erbium, and "ytterbium" is 56 approximately?

In accordance with Ostwald's Rule's the value of  $\lambda_{1021} = \lambda_{-1}$  should be about 30 for the chlorides and intrates and about 60 for the sulphates of the rare oaith elements are tervalent, the corresponding values for bivalent metals being 20 and 10 respectively. It is evident that the experimental evidence is, on the whole, decidedly favourable to the tervalency of the metals?

<sup>1</sup> Matignon, Ann Chim Phys., 1906, [viii.], 8, 274 · Nd), 403 (Sm., 444 · (Yb)), other data due to Baoult, ibid., 1885, [vi.], 4, 411.

2 W. Biltz, Annalen, 1904, 331, 334

3 The formulae are written on the assumption that the rare earth elements are tervalent, but the minerical data, being equivalent conductivities, are independent of this assumption. The dilution ris expressed in hires per grain equivalent, as usual.

4 Ley, Zatsch. physikal Chim., 1899, 30, 193.

5 Anfrecht, Inanquial Dissertation (Berlin, 1904), see Abegg, Handbuch der anorganischen Chemie (Leiping, 1906), vol. 3, pt. 1.

<sup>\*</sup> Antrecht, Inauqueal Dissentation (Berlin, 1904), see Abegg, Handbuch der anorganischen Chemie (Leipzig, 1906), vol. 3, pt. 1.

\* Muthmain, Her, 1898, 31, 1829

† Brauner and Svage eited in Abegg, opus ed., p. 170

For other conductivity measurements, see Jones and Allen, Amer. Chem. J., 1898, 18, 321. Jones and Reese, ibid., 1898, 20, 600; Noyes and Johnston, J. Amer. Chem. Suc., 1909, 31, 987

\* See Vol. I. p. 150

† That the rare earth elements are tervalent is also shown by the behaviour of their salts in coagulating arsenious sulphide hydrosol (Freundlich and Schucht, Zeitsch. physikal. Chem., 1912, 30, 564).

(vi.) Composition of Salts.—The assumption that the rare earth elements are triads often leads to formula considerably simpler than those necessitated by the assumption that they are diads. This was pointed out by Cleve and Hogland in discussing the compounds of yttimm and "cibium". The following alternative formulæ of a number of yttimm salts may be cited.—

The manner in which  $Y_i$  enters into the diad formulae of the double salts is very curious and significant

Similar results hold for the salts of each of the rare earth metals that has been studied in any detail. The compounds that cerous neodynamic, and samarium chlorides form with animonia may be cited as further examples of the simplicity of the triad formula.

The triad formula are here clearly more convenient than the diad

Further than this, the double and complex salts of the rare earth elements are often different in type from those of the known dual elements. Consider, for instance, the iddinaplatinates. Those of the known unit and breadent metals are normal, is the chlorine of the platine chloride is twice the chlorine of the metal chloride, i.g. 2KCl PCCl, and MgCl, PCCl, 6H,O. The salts of metals known to be trivalent are not, however, of the bornal type, but are basic, and the salts of the rare earth elements are similar in composition to them, thus

The indium and yttrium salts are anomalous, being 21nCl\_5PtCl\_36H\_O and 4VCl\_5PtCl\_52H\_O respectively, but neither is of the normal type. From the composition of the chloroplatmates, then, the rare earth elements must

<sup>&</sup>lt;sup>1</sup> Cleve and Hoglund, Bull Soc chim., 1872, [n.], 18, 193, 289. See also Cleve, did., 1874, [n.], 21, 196, 246, 344, John, ibid., 1874, [n.], 21, 533.

be regarded as tervalent, and the same conclusion may be drawn from a study of the chloroplatinites and the selecutes 1

(vn.) The Periodic Classification — It has been already stated that Mendeléess was the first to reject the idea of the bivalency of the rare earth elements, and the formula he proposed for the oxides of these elements have been given (see p. 231). His method of incorporating the rare earth elements into the Periodic Table was as follows —

| Sı |    | Y  |    | Zı |    | Nb |    |
|----|----|----|----|----|----|----|----|
| Ва | Cd | Dı | In | Ce | Sn |    | Sb |
|    |    | Er | -  | La | i  | Ta | -  |

With regard to these positions, Mendelecti was convinced of the accuracy of his views concerning cerium, the more so as he showed experimentally that cerium thereby followed. Dulong and Petits Law, but although he was satisfied with the position he assigned to yttrum, he was doubtful over lanthanum, didymnum, and erbinin. The positions assigned to these elements by Mendelecti in his table need not, however, be further discussed, since the "didymnum" and "erbinin" of 1872 are now known to be complex mixtures.

At the present time the problem is to place not five, but fifteen, rare earth elements into the Periodic Table. Inspection of that table will show that in seeking positions for the rare earth elements only two vacant places occur for elements of atomic weight less than that of barinin, viz one between strontium (87:63) and zincomini (90:6), and the other between molybdemini (96:0) and ruthenium (101:7). Between barium (137:37) and tantalum (181:0) there are many vacant places. The atomic weights of the rare earth elements are as follows, according to whether they are diads or triads:—

|                                                           |              |                         |                                                              |                         |                         |                                                       |       |                | ; |
|-----------------------------------------------------------|--------------|-------------------------|--------------------------------------------------------------|-------------------------|-------------------------|-------------------------------------------------------|-------|----------------|---|
| Element                                                   | Diad         | l'uad                   | Element                                                      | Diad                    | Triad                   | Element                                               | Drad  | Triad.         |   |
|                                                           | ŀ            | İ                       | -                                                            | ļ                       |                         |                                                       | '     |                | ĺ |
| Yttmum<br>Lanthanum<br>Cemum<br>Praseodymmum<br>Neodymmum | 93 5<br>93 9 | 189.0<br>140.3<br>140.9 | Sam trium<br>Europium<br>Gadolinium<br>Terbium<br>Dysprosium | 101 3<br>101 9<br>106 1 | 152 0<br>157 3<br>159 2 | Holmum<br>Erbium<br>Thulium<br>V tterbium<br>Lutecium | 115 7 | 167 7<br>168 5 |   |
|                                                           |              |                         |                                                              | 1                       | 1 1                     |                                                       |       | ا _ا           | ı |

Assuming that the rare earth elements are diads, it is seen at once that (i.) there is no place in the Periodic Table for yttrium, (ii) three elements

<sup>1</sup> Nilson, Ber., 1876, 9, 1056 (chloreplatmates), Ber., 1875, 9, 1112, or J. pickl Chem., 1877, [n.], 15, 260 (chloreplatmates), Ber., 1875, 8, 655, Bull Soc. chim., 1875, [n.], 23, 494; or Nova Acta Soc. Upada, 1875, [n.], 9, No. 7 (scientes). See also Cleve and Hoglund, Bull. Soc. chim., 1872, [n.], 18, 193, 289, Cleve, ibid., 1874, [n.] 21, 196, 246, 344.

8 See p. 232.

compete for the one vacant place between molybdenum and rathennum, (iii.) eleven other elements rescuble yttrium in being without places, and (iv.) the spaces between bariam and fantalim still remain inoccupied. On the other hand, assuming the elements to be triads, it is observed that (i.) yttrium occupies the vacant space between strontium and zirconnum, (ii) the space between molybdenum and rathernum remains vacint, and (iii) fourteen elements, the atomic weights of which he between those of barium and tantalium, are available for filling the vacance—between the positions of those two elements.

On the assumption of the tervalency of the rare earth elements, then, it possible to place all these elements into vacant spaces. From the chemical point of view, however, the positions that the rare earth elements then occupy are most unsatisfactory, except in the cases of yttrium, lanthamum, and cerium. Lanthamum and cerium follow directly after barium, for there are good reasons for regarding cerium as a member of Group IV in the table. Hence, scandinm, yttrium, and lanthamim must be grouped together as constituting the even subgroup of Group III. and this view is in very fair accordance with their properties. There still remain, however, twelve more elements, and it is impossible to arrange them in positions which range from Group I to Group VIII, to assume that each is torvalent in its salts, and still accept the truth of the Periodic Law. Clearly such assumptions are mutually contradictory. So far, therefore, as the rare earth elements are concerned, Mendeleeff's classification is defective, and cannot be legitimately utilised in arguing for or against the tervalency of the rare earth elements.

(vm.) The Case of Cerium.—The element cerium differs from the other rate earth elements in that it forms two basic oxides and two series of salts. Reterred to unit weight of cerium, the oxygen in the lower oxide is to that in the Ligher oxide as 3 is to 4. From the point of view or simplicity, then, the oxides are best formulated as Ce<sub>2</sub>O and CeO<sub>2</sub> ceriam being regarded as both trial and tetrad. If, however, cerimin in cerous salts is regarded as a dual, it follows that the higher oxide is Ce O<sub>2</sub> quite a unique type of basic oxide, and dual the formulae of the cerie salts are very complicated.

There is, moreover, strong evidence in favour of the first of the two preceding views, and for classifying cerimin with titanium, zneomium, and thorium as a group of tetral elements. Since the ovidence is fully discussed in a subsequent volume, it is immecessary to repeat it here, it is only necessary to point out that the fact that evidence independent of the preceding shows cerimin to be terralent in croins salts is a strong argument in favour of the tervalency of the other rare earth elements.

(ix.) Summary - The evidence bearing upon the question of the valencies and atomic weights of the raie earth elements has now been discussed, from various different points of view. Whatever the value that may be assigned to the crystallographic evidence and the appeal to the Periodic Law, the balance of evidence indicates that the raie earth elements are tervalent in their salts. This view, which is accepted by nearly all chemists, is the one adopted in this book.

Practical Methods of Atomic Weight Measurement.—The chief methods that have been employed in the determination of the atomic weights of the rare earth elements are briefly outlined in the following paragraphs.

(1.) Sulphate Methods. - (a) Synthetical. A weighed amount of the pure sesquioxide, prepared by the ignition of the pure oxalate, is dissolved in hydrochloric acid in a weighed platinum crucible, a slight excess of sulphuric acid added, and the solution evaporated as far as possible on the water-bath. The evaporation is then continued at a higher temperature until the excess of sulphurie acid has been removed and the residual anhydrous sulphate brought to constant weight at about  $500^\circ$  . If the weights of oxide and sulphato be aand b respectively, and the required atomic weight be X, then

This method, originated (in a slightly different form) by Bahi and Bunsen in 1866, has been used more than any other for determining the atomic weights of the rare earth elements.1 Although, however, the procedure is simple, the method is certainly not the best that can be employed for the atomic weight determinations. Owing to the extremely hygroscopic nature of the anhydrous sulphates, great care must be taken to exclude moisture from the salts until they have been weighed. The chief difficulty, however, consists in obtaining the sulphates dry and neutral, since water and sulphuric acid are retained if the final temperature of heating is too low, and decomposition of the sulphates begins if the final temperature is too high. In the former case, the sulphates dissolve in water, forming clear solutions which react acid towards methylorange, in the latter, the sulphates do not dissolve to a clear solution, The precise final temperature to employ probably requires in each individual case a special investigation for its determination. In many cases, the boiling-point of sulphur, which is probably too low, has been chosen. Wild recommends 450'-500', and other chemists have adopted 500". According to Brauner, it is always necessary to dissolve the residual sulphate is water, titrate with N, 10 sodium hydroxide till the solution is neutral, and apply the appropriate correction. In view of the results obtained by Jones and by Holmberg, however, it is difficult to behave that the preparation of a dry,

neutral sulphate cannot be accomplished  $^2$  (b) Analytical—in this process the octahydrated sulphate,  $X_4(SO_4)_3$   $SH_2O_4$  is chosen as starting material—The sample for analysis may generally be prepared by dissolving a gram or two of oxide in hydrochloric acid, evaporating nearly to dryness, diluting to fifty cubic centimetres, adding a slight excess of sulphine acid, and then quickly diluting with a litre of alcohol. The precipitated sulphate is well washed with alcohol, dired at 110°, dissolved in water, and crystallised at the temperaturo of the water bath The crystals are separated from the mother liquor, dired in the an, powdered in an agate mortar, and dried in a desiceator. A gram or two of the sulphate

<sup>1</sup> For further details of the method see Babi and Buosen, Innalen 1866, 137, 1. Bailey, 1 For further details of the method see Babr and Bunsen, Innat. in 1865, 137, 1, Bailey, Trans Chm. See, 1885, 51, 683, Kines, Zeitsk, amay Chm., 1893, 3, 14, Utham, Ann Chim Phys., 1900, [81], 19, 184, Jones, Amer. Chem. J., 1902, 28, 23, Chem. News, 1903, 38, 13, Zeitsk amarg. Chem., 1903, 36, 92, Biauner and Patheck, Trans. Chem. Soc., 1902, 81, 1243, Biauner, Lutsch amay Chem., 1903, 33, 317, Whl, John 1904, 38, 191; Bull, whid., 1905, 47, 164, Holmberg, whid., 1907, 53, 124, R. J. Meyer and Wuormen, whid., 1913, 80, 7, Egan and Balke, J. Amer. Chem. Soc., 1913, 35, 365, 2 It is possible, as Meyer and Wuormen have pounted out, that the duboulites may have usually ansen from the fact that the oxide has been ignited in platinum and contaminated with a trace of that element, and hence that the rale earth sulphate subsequently formed has been sightly contaminated with a trace of that element, and hence that the rale earth sulphate subsequently formed has been sightly contaminated with a trace of that element, and hence that the rale earth sulphate subsequently formed has been sightly contaminated with a trace of that element.

has been slightly contaminated with platinic sulphate, of Jones, loc. cit

is weighed into a small platinum crucible, and heated to 350° till the water of crystallisation has been expelled. The loss of weight is then determined. Finally, the crucible and contents are placed in a larger crucible (which it is prevented from touching by means of a little ignited rare earth packing) and heated almost to a white heat until the sulphate has been completely decomposed. The weight of the residied oxide is then determined

By this method, in addition to the oride sulphate ratio (L), values for the atomic weight may be calculated from two other ratios, viz

$$X_i(80_4)_i 81I_i 0 - X_i(80_4)_3$$
 (11)

and

$$X_{i}(SO_{i})_{3}SH_{i}O_{i} = X_{i}O_{3}$$
 (111)

It should be noted that with cerum, the final residue is the diovide With praseodynaum and terbium, the final residues are peroxides of some what indefinite composition. They may, however, be reduced to sesqui oxides in a stream of hydrogen.

This method has been employed very successfully by Urbain, who prefers the values deduced from ratio (111) to the others. The oracle sulphate into as determined by the analytical method avoids the difficulty, encountered in the synthetic method, or driving off excess of acid, but it suiters from another defect. The anhydrous sulphate obtained is extremely porous, and absorbs gases, so much so that it is impossible, for just once, to weigh anhydrous gadolimina sulphate in air with any reasonable degree of accuracy

The foregoing method may be applied to very small quantities of material, the weighing being neide on a torsion interobabance.

According to 15 omer, the analytical method is bouted in its scope, since it is impossible to decompose completely by ignition the sulph ite of landhamin and the other sulphates derived from the strongly basic cortles. The analytical method is therefore best sinted to the sidph descot the elements of the terbinic and cibring groups. As an illustration of the retine of the results it has afforded, the following data referring to the atomic weight of emopium,

| No. of   | 1           | Weight of                      |                   | ' Atomo W | right of Eu | from Rati |
|----------|-------------|--------------------------------|-------------------|-----------|-------------|-----------|
| å e tion | Tu/80/\SH20 | $\mathrm{Eu}/\mathrm{SO}_4)_t$ | Ea_O <sub>3</sub> | 1         | 11          | 111       |
| 15       | 1.7787      | 1 4303                         | 0 > 201           | 151-91    | 151 74      | 151.83    |
| 16       | 2 1785      | 1 9935                         | 1 1515            | 1 (1 95   | 172 10      | 151 98    |
| 17       | 2 1177      | 1 9149                         | 1.1554            | 151 76    | 152 33      | 151.88    |
| 18       | 211801      | 1 0068                         | 1 1470            | 152 04    | 111.80      | 151 98    |
| 19       | 2 2988      | 1 5185                         | 1 0990            | 152 03    | 151 196     | 152 01    |
| -        |             |                                | 1                 |           |             | ٠         |
|          |             | Me                             | l)                | 151 94    | 151 98      | 151:94    |

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Urham, J. Chim, phys., 1966. 4, 52
 Birll, he cit. of Nerrest art Rosa (Ed., Ber., 1993, 36, 2086).
 Urham and Lacombe, Com. t. read., 1994, 138, 627. J. Urham, J. Chim. phys., 1996, 118. The autecodem data are O=16, H=1,00762, S=32,005. The numbers attached 4, 118 The antecedent data are 0=16, n=1 corresponds to the fractions have the same meanings as those on p  $-34^{\circ}$ 

(ii.) The Oxalate Method - This method is more suited to the rapid determination of approximate results than to the determination of exact atomic weight values. The precipitated oxalate is either an-dried or dried at 120°-130° and ground in an agate mortar to ensure its homogeneity. In one portion the percentage of rare earth oxide is determined by ignition,1 and in another the oxalate radicle is estimated by titration with a standard solution of potassium permanganate, or by an ordinary combustion analysis. From the ratio X<sub>2</sub>O<sub>3</sub> 3C<sub>2</sub>O<sub>3</sub> the atomic weight is easily calculated <sup>2</sup> The oxalate should be prepared by precipitation with oxalic acid from a slightly acid solution, if precipitated by a soluble oxalate from a neutral or alkaline solution, the oxalates of the alkali metals and ammonium are co-precipitated to a very considerable extent.8

(m) The Halide Methods —(a) The analysis of the pure, anhydrous chloride is an operation comparatively easy to effect, and when properly performed always yields very reliable results. The preparation of the pure, anhydrous chlorides, however, is a difficult task, so that this method has unfortunately been greatly neglected, 4 analysis of the bromides, which should give equally

good results, have not yet been attempted

(b) The determination of the ratio X2O, 2XCl, may be effected by dissolving the oxide, contained in a quartz flask, in hydrochloric acid, evaporating to dryness in a stream of hydrogen chloride, and dehydrating

and fusing the residual chloride in a current of the same gas

(iv.) Other Methods - A number of useful, rapid volumetric processes, which, however, yield only approximate results, have been described " A convenient procedure is to dissolve about 0 1500 gram of the pure oxide in 50 cc of standard sulphuric acid (approximately seminormal), add about 8 c.c of N/5 potassium oxalate, and heat to the boiling-point, cool and titrate the excess of sulphurie acid with sodium hydroxide, using phenolphthalem as indicator

Experimental Results.-The experimental results which seem to be the most reliable are contained in the accompanying table, to which a fairly complete bibliography is appended. So far as the atomic weights are concorned, many of the papers referred to in the bibliography have only an historical interest.

<sup>1</sup> It is difficult to effect this determination without a trace of oxide being mechanically

It is difficult to effect this determination without a trace of oxide being mechanically carried awily by escaping gases.
 Stolba Rev bokin Ges Wiss., 1878., Johnesber., 1878., p. 1959., 1879. p. 1014., 1882., p. 1286., Gibbs., Proc. Amer. Acad., 1893., 28, 261., Amer. Chem. J., 1893., 15, 516.
 Biaumer, Chem., New., 1895., 71, 233., Branner and Paxhéek, Joe. at.; Brauner and Batek, Zettsch. amorg. Chem., 1903., 34, 103., Branner, Ind., 1903., 34, 207., Browning. Amer. J. Ser., 1899., [vr.], 8, 451., Drushel, Ibid., 1907., [vr.], 24, 197.
 Baxter and Griffin, J. Amer. Chem. Soc., 1906., 28, 1681; Baxter and Daudt, Ibid., 1908., 30, 563.
 For determinations by this method, see Robinson, Proc. Roy. Soc., 1884., 37, 150., Chem. New., 1884., 50, 251., Baxter and Chapin, J. Amer. Chem. Soc., 1911., 33, 1., Baxter and Stewart, Ibid. 1915., 37, 516.
 Egan and Balke, J. Amer. Chem. Soc., 1913., 35, 365.
 See Kruss and Louce, Zettsch. amorg. Chem., 1883., 4, 161; Wild. Ibid., 1904., 38, 191., Fert and Prablylla, Ibid., 1905., 43, 202; 1906., 50, 249., Holden and James, J. Amer. Chem. Soc., 1914., 36, 638.

# ATOMIC WEIGHTS OF THE RARE EARTH ELEMENTS.1

| Date                                 | Authority.                                                                    | Ratio Measured.                                                                                                                                                                                                           | Atomic Weight Value (1916).                               |
|--------------------------------------|-------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------|
|                                      |                                                                               | Cirium 2                                                                                                                                                                                                                  |                                                           |
| 1884<br>1885<br>1897<br>1903<br>1903 | Robinson<br>Brauner<br>Wytouboff and Verneuil<br>Branner and Batck<br>Brauner | $ \begin{array}{l} \left[ \begin{array}{ccc} CCI & \exists Ag \\ (e_2/SO_4)_1 & 2CeO_2 \\ Ce_2/SO_4 & 8H_2O & Ce_2/SO_4 \end{array} \right] & 2CeO_2 \\ Ce_2/SO_4)_3 & 2CeO_2 \\ Ce_2/SO_4 & 8H_2O & 2CeO_2 \end{array} $ | 140 14* 140 25<br>140 22*<br>149 35<br>140 30*<br>140 20* |
|                                      |                                                                               | Dysprosium                                                                                                                                                                                                                |                                                           |
| 1906                                 | Urbain and Demenitions                                                        | Dy <sub>2</sub> (8O <sub>4</sub> ) <sub>3</sub> 8H <sub>2</sub> O Dy <sub>3</sub> O <sub>3</sub>                                                                                                                          | 162 55   162 5                                            |
|                                      |                                                                               | Евви м 4                                                                                                                                                                                                                  |                                                           |
| 1908  <br>1910                       | Hotmann and Burger<br>Hofmann                                                 | $\begin{array}{c} \operatorname{El}_2\mathrm{O}_3:\operatorname{Er}_2(\mathrm{SO}_4)_1 \\ \operatorname{El}_2\mathrm{O}_3 - \operatorname{El}_2 \operatorname{SO}_4\rangle_1 \end{array}$                                 | 167°40 ± 167°7<br>  167°68                                |
|                                      |                                                                               | Епкори м.                                                                                                                                                                                                                 | 1                                                         |
| 1908  <br>1908                       | Urbain and Lacombe<br>Jantsch                                                 | $\begin{array}{lll} \operatorname{En}_f(SO_4)_2 & SH & O & \operatorname{En}_f(SO_4)_4 & \operatorname{En}_2O_4 \\ \operatorname{En}_4 & SO_4)_4 & SH_2O & \operatorname{En}_2O_3 \end{array}$                            | 151 94<br>162 05   152 0                                  |

<sup>&</sup>lt;sup>4</sup> With one or two exceptions, the results given in the accompanying table have been computed from the data given in Carke, A Readendation of the Alonce Weintle (\*\*Smithsuman Miscellam ous Collections, \*\*vol. 54, No. 3, 1910\*), using the following anter-dent data \*\*\*\*

In the few cases indicated by asterisks the weights have been reduced to vacuum standard in calculating the results.

The cather exp innenters never freed their certa completely from "didyima"

Dysprosia was not isolated till 1906.

Pure ciba has not yet been prepared
 Europia was not isolated in the pure state till 1904

Cerum — Beringer, Annalen, 1842, 42, 134, Hermann, J. prakt, Chem., 1813, 30, 185; Marignae, Arch. Sci. phys. nat., 1818, [1], 8, 273, h. perulf, Annalen, 1853, 87, 12. Bunsen and Jegel, ibid., 1855, 105, 45, Rammelsherg. Popy. Annalen, 1859, 108, 44; Wolf, Amer. J. Sci., 1868, [10], 46, 63, Wing, ibid., 1870, [1], 49, 378; Buhring, J. prakt. Chem., 1875, 120, 222, Robinson, Proc. Roy. Soc., 1884, 37, 150, Chem. Neus, 1881, 50, 251, 1886, 54, 229; Bhanner, Trans. Chem. Soc., 1885, 47, 879, Schotzenberger, Compd. road., 1895, 120, 663, 962, 1143. Brauner, Chem. Neus, 1895, 71, 283, Wyroubolf and Veincul. Ball. Soc., Chim., 1897, [11], 17, 679, Compd. road., 1895, 71, 283, Wyroubolf and Veincul. Ball. Soc., Chim., 1897, [11], 17, 679, Compd. road., 1897, 124, 1300, Ann. Chem. Phys., 1907, (viii.), 9, 349, Kolle, Inaugural Dissertation Zurich, 1899); Brauner and Battch. Zedsch. anary. Chem., 1903, 34, 207.

Dysprosium.—Urbain and Deminitroux, Compt. rend., 1906, 143, 598

Erbium.—Cleve, K. Svenska Vet. Akad. Handl., 1880, No. 7; Compt. rend., 1880, 91, 381; Brill, Zeitsch. anary. Chem., 1905, 47, 464; Hofmann and Buiger, Rev., 1908, 41, 308; Hofmann, Rev., 1910, 43, 2635

Europum.—Urbain and Lacombe, Compt. rend., 1904, 138, 627, Urbain, J. Chem., phys., 1906, 4, 118, Jantsch, Compt. rend., 1908, 146, 473, cf. Demais, Jubid., 1900, 130, 1469; 1901, 132, 1484.

 $O=16\ 000$  ,  $S=32\ 065$  ,  $H=1\ 00762$  , Ag  $-107\ 880$  ,  $Cl=35\ ^{157}$  ,  $C=12\ 003$ 

ATOMIC WEIGHTS OF THE RARE EARTH ELEMENTS-continued.

| Date                 | Authority.                                                      | Ratio Measured.                                                                                                                                                                                                                                                                                                                              | Atomic Unitional Value (1916).           |
|----------------------|-----------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------|
|                      |                                                                 | GADOLINIUM 1                                                                                                                                                                                                                                                                                                                                 |                                          |
| 1905   I             | Irbarn                                                          | $[\operatorname{Gd}_2(\operatorname{SO}_4)_1 \operatorname{SH}_2\operatorname{O} \operatorname{Gd}_2\operatorname{O}_3$                                                                                                                                                                                                                      | 157 21   157 3                           |
|                      |                                                                 | Подмичм 2                                                                                                                                                                                                                                                                                                                                    |                                          |
| 1911   I             | folmberg                                                        | Ho <sub>3</sub> O   Ho <sub>2</sub> (SO <sub>4</sub> ) <sub>1</sub>                                                                                                                                                                                                                                                                          | 163 45   163 5                           |
|                      |                                                                 | LANTHANUM                                                                                                                                                                                                                                                                                                                                    |                                          |
| 1902                 | iones<br>Braunci and Pavliček<br>Jones                          | La <sub>1</sub> O <sub>1</sub>   La <sub>2</sub> (SO <sub>4</sub> ) <sub>1</sub>   La <sub>2</sub> O   La <sub>2</sub> (SO <sub>4</sub> ) <sub>1</sub>   La <sub>2</sub> O <sub>3</sub>   3C <sub>2</sub> O <sub>3</sub>   La <sub>2</sub> O <sub>3</sub>   La <sub>2</sub> O <sub>3</sub>   La <sub>2</sub> (SO <sub>4</sub> ) <sub>2</sub> | 139.05*<br>139.05*<br>139.09*<br>138.76* |
|                      |                                                                 | Laterum (or cassiopeium) 3                                                                                                                                                                                                                                                                                                                   |                                          |
| 1907 8 U<br>1913   A | Jibain<br>Auer von Welsbach                                     | $\begin{array}{c} \mathrm{Lm}_{3}(\mathrm{SO}_{4})_{3} \ 8\mathrm{H}_{2}\mathrm{O} : \mathrm{Lu}_{2}\mathrm{O} \\ \mathrm{Lu}_{2}(\mathrm{SO}_{4})_{3} \ 8\mathrm{H}_{2}\mathrm{O} & \mathrm{Lu}_{2}\mathrm{O} \end{array}$                                                                                                                  | 171 175 0<br>175 00                      |
|                      |                                                                 | Ni odymium,                                                                                                                                                                                                                                                                                                                                  |                                          |
| 1907                 | Auer von Welsbach<br>Holmberg<br>Baxter and Chapin <sup>4</sup> | Nd <sub>2</sub> O <sub>1</sub>   Nd <sub>2</sub> (SO <sub>4</sub> ) <sub>1</sub>   Nd <sub>2</sub> O <sub>4</sub>   Nd <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub>   NdCl <sub>1</sub>   3AgCl   NdCl <sub>1</sub>   3Ag                                                                                                                    | 141.55 111.3<br>111.10<br>1.11.27*       |

 $<sup>^{1}</sup>$  Earlier determinations are most probably affected by presence of europia in the gadolima used

Holma was not isolated until 1911

It is doubtful whether pure interia has yet been obtained
 New analyses of NdCl<sub>1</sub> by Bayter, Whitcomb, Stewart and Chapin confirm the results of Bayter and Chapin (J. Amer. Chem. Soc., 1916, 38, 302)

Gadolinium. - Mangiac. Aan. Chim Phys., 1880, [v.], 20, 535. Levoq de Borsbandian, Compt. rend., 1890, 111, 109; Bottendorf, Annalea. 1892. 270, 578. Benedicks. Zeitsch annay Chem., 1800, 22, 393. Mar., thid., 1904, 38, 128. Demaid., Compt. rend., 1800, 131, 343. Urbain, ibid., 1905, 140, 583. J. Chim. phys., 1906, 4, 326.
Holmium. - Holmberg. Arki. Kim. Min. Giad., 1911, 4, No. 2; Zeitsch anory Chem., 1912, 71, 226.
Lanthanum. - Mosander, Pogg. Annalea, 1843, 60, 297. Rammelsbeig, ibid., 1812, 55, 65; Mangiac. Arch. Sci. phys. nat., 1849, 11, 21. Ann. Chim. Phus., 1819, [iii.], 27, 209; Holzmann, J. patt. Chem., 1868, 704, 174. Etc., J. nacrah. Zeitsch., 1871, 6, 396. Mangiac. Arch. Sci. phys. nat., 1883, 61, 4174. Etc., J. nacrah. Zeitsch., 1871, 6, 306. Mangiac. Arch. Sci. phys. nat., 1873, 46, 215. Ann. Chim. Phus., 1873, [iv.], 30, 68; Cleve, Bihang K. Scienku Vet. Alkad. Handl., 1874, 2, No. 7, Bull. No. chim., 1874, [ii.], 21, 196, 1883, [ii.], 39, 151. Bianner, Monalsh., 1882, 3, 1, 456; Trons. Chem. Noc., 1882, 41, 68. Bei., 1891, 24, 1328. Bettendorf, Annalea, 1890, 256, 159. Gibbs., Proc. Amer. Acad., 1893. 18, 260, Schntzenberger, Compt., iend., 1895, 120, 1113. Bodman, Bihang K. Scienka Vet.-Akad. Handl., 1901, 26, II., No. 3; Brauner and Pachick, Trans. Chem. Soc., 1902, 81, 1243. Bhauner, Zeitsch anorg. Chem., 1903, 33, 317. Jones., imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem. J., 1902, 28, 23, Zeitsch anorg. Chem., 1903, 33, 317. Jones, imer. Chem

### ATOMIC WEIGHTS OF THE RARE EARTH ELEMENTS-continued.

| Date Authority.                                                                                        | Ratio Measured                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | Atomic national<br>Weight : Value<br>(1910)                                              |
|--------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------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| 1                                                                                                      | Praskodymu m                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |                                                                                          |
| 1898 John 9<br>1898 Von Scheele<br>1901 Brauner V<br>1993 Auer von Welsbach<br>1915 Baxter and Stewart | Pr <sub>2</sub> O <sub>4</sub>   Pr <sub>4</sub> (SO <sub>4</sub> )   Pr <sub>4</sub> O <sub>5</sub>   Pr <sub>4</sub> (SO <sub>4</sub> )   Pr <sub>4</sub> O <sub>5</sub>   Pr <sub>4</sub> (SO <sub>4</sub> )   Pr <sub>4</sub> O <sub>5</sub>   Pr <sub>4</sub> (SO <sub>4</sub> )   Pr <sub>4</sub> O <sub>5</sub>   Pr <sub>4</sub> O <sub>4</sub>   Pr <sub>4</sub> O <sub>5</sub>   Pr <sub>4</sub> O <sub>4</sub>   Pr <sub>4</sub> O <sub></sub> | 140 47   140 9<br>110 54  <br>110 97*  <br>110 96*  <br>140 57  <br>140 93*  <br>110 92* |
|                                                                                                        | SAMABIUM 1                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |                                                                                          |
| 1904   Urbain and Lacombe                                                                              | Sm <sub>2</sub> (SO <sub>4</sub> ) 8H <sub>2</sub> O   Sm <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> , Sm <sub>2</sub> O <sub>4</sub>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 150 16   150 4                                                                           |
|                                                                                                        | Tarbium 2                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         |                                                                                          |
| 1906 . Urbain                                                                                          | Tb <sub>2</sub> SO <sub>4</sub> \SH <sub>2</sub> O   Tb <sub>2</sub> (SO <sub>4</sub> ) <sub>4</sub>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 159 20   159 2                                                                           |
|                                                                                                        | THI LILY                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |                                                                                          |
| 1880 Cleve<br>1905 Urban                                                                               | 1                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 170 7<br>  168 5   168 5                                                                 |
| Yerren                                                                                                 | rn (neagterlaum or aldebaranium)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |                                                                                          |
| 1913 , Aner von Welsbach<br>1914 Blumonfeld and Urbain                                                 | $\frac{\text{V} \text{b}_2(\text{SO}_4)_3 \text{ SH}_3\text{O} - \text{V} \text{b}_2\text{O}_4}{\text{V} \text{b}_2(\text{SO}_4)_6 \text{ SH}_2\text{O} - \text{V} \text{b}_2\text{O}_4}$                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 173 0<br>173 74                                                                          |

- Samaria was not obtained free from neodymna and emopia until 1900.

  - f crbia was not isolated until 1906
     A reliable determination is still wanted

Biss B a.a., 1903, 112 H. B., 1037, Brauner, Proc. Clem. Soc., 1898, 14, 72, 1901, 17, no., Abeggs Handland, vol. 3 part 1, p. 276, Bondonard, Compt. rend., 1898, 126, 900; Jones, Inar. Chem. J., 1898, 20, 315, Zeitsch. nangr. Chem., 1899, 19, 329. Brill, And., 1905, 47, 104, Holmberg, ibad, 1907, 53, 124, Fert and Pribyllis, ibad, 1906, 50, 259; Baxter and Chapin, J., Inar. Chem. Soc., 1911, 33, 1.

Praseodymium — Augr. von Welsbich, Ios. et al. Braimer, Proc. Chem. Soc., 1898, 14, 70, 1801, 17, 65, Abegg's Handbuch, vol. 3, part 1, p. 203, Jones, Amer. Chem. J., 1898, 20, 315, von Scheele, Zeitsch. nang. Chem., 1898, 17, 310, Fert and Pribyllia, and., 1906, 50, 258. Baxter and Stewart, J. Amer. Chem. Soc., 1915, 37, 516.

Samarium. Maniquae, Irch. Soc., phys. and., 1890, 101, 3, 435, Ann. Chem. Phys., 1880, [v.], 20, 535, Brainer, Trans. Chem. Soc., 1883, 43, 278, Cleve, ibid., 1883, 43, 362, Bettendorf, Annalen, 1891, 263, 164, Lacoqde Borshandian, Compt. rend., 1892, 114, 575. Demagay, ibid., 1896, 122, 728, 1900, 130, 1919, 1185, 1409, 1901, 132, 1484, Muthmann and Ways. Annalen, 1901, 331, 16, Urbain and Laconbe, Compt. rend., 1904, 138, 1166, Urbain, J. Chem. Phys., 1900, 4, 120, Matignon, Compt. rend., 1905, 141, 1230; Fert and Pribylla, Zeitsch. anoig. Chem., 1905, 50, 259.

Terbium.—Urbain, Compt. cend., 1906, 142, 957, cf. Fert, Zeitsch. anoig. Chem., 1905, 43, 280, Lecoq. de Borshandian, Compt. rend., 1880, 91, 50, A81, Compt. rend., 1914, 159, 323. Sec. also the reference given under luterium. For the "atomic weight" of old "ytterbium," see Nilson, Compt. rend., 1880, 91, 50, Astrid Cleve, Zeitsch. anoig. Chem., 1902, 32, 129, Brill, 1904, 1905, 47, 364, Brainer, Abegg's Handbuch, vol. 3, part., p. 335, G. and., 50, 261.

50, 261.

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#### ATOMIC WEIGHTS OF THE RARE EARTH ELEMENTS-continued.

| Date         | Authority                                                                 | Ratio Measured                                                                                                                                                                                                                                                                                                         | Atomic<br>Weight                  | Inter-<br>national<br>Value<br>(1916) |
|--------------|---------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------|---------------------------------------|
| 1895<br>1913 | Cleve<br>Jones<br>R. J. Meyer<br>Wuorinen and Weinheber<br>Egan and Balke | $\begin{array}{c} {\rm Yr}_{\rm IRIUM}, \\ {\rm Y}_2{\rm O}_3 - {\rm Y}_2({\rm SO}_4)_3 \\ {\rm Y}_2{\rm O}_1 - {\rm Y}_2({\rm SO}_4)_3 \\ {\rm Y}_2{\rm O}_1 - {\rm Y}_2^{\prime}{\rm SO}_4)_3 \\ {\rm Y}_2^{\prime}{\rm O}_3 - {\rm Y}_2^{\prime}{\rm O}_4 \\ {\rm Y}_2^{\prime}{\rm O}_3 - {\rm 2YO}_3 \end{array}$ | 89·12<br>88·97<br>88·70<br>90·101 | 88.7                                  |

<sup>&</sup>lt;sup>1</sup> Preliminary results, and admittedly a little high.

Yttrium — Beilin, Finhandl Skand Nat Kjob., 1860, p. 452, Popp. Annalen, 1861, 131, 179. Delafontame, that, 1865, 134, 108. Arch Nev phus nat, 1866, [u.], 25, 119. Bahr and Binsen, Annalen, 1866, 137, 1, Cleve and Hogland, Bihang K. Nevaska Vettakad Hemill., 1873, No. 8. Cleve, that, 1882, No. 9. Bull. Soc. chim., 1883, 101, 39, 120, 287. Compit rend., 1882, 95, 1225, Jones, Amer Chem. J., 1895, 14, 151, Chem. News, 1895, 71, 305, Mathimann and Bohm, Ber., 1900, 33, 42; G. and E. Union, Loc. cit. Brill, Ion. cit. Feet and Privibile, Zeitsch. anorg Chem., 1906, 262, R. J. Meyer and Wuotinen, 4bid., 1913, 80, 7, R. J. Meyer and Weinheber, Ber., 1913, 46, 2672, Egan and Balke, J. Amer Chem. Soc., 1613, 35, 365.

It is difficult to form any precise estimates of the possible errors attaching to the atomic weight values chosen for the raio carth elements by the international committee. It is possible that the values ascribed to ceruin, prascodymium, neodymium, dysprosium, europium, gadolimium, terbium, samarium, yttrium, and yttribium do not err by more than one- or two-tenths of a unit from the correct values, and that in three or four cases the error does not exceed one or two units in the second decimal, it is possibly piemature to express the value for ceruin to two places of decimals. In the case of lainthanium there is a discrepancy of nearly three-tenths of a unit between the values of Jones (138-73) and Braunei (193-05), and the evidence in favour of the lower value is at least as convincing as that for the higher Considerable uncertainty also attaches to the values for erbium, thulium, and lutecium, while additional determinations are required to check the value ascribed to holimium. No values have yet been published for the atomic weight of celtium <sup>1</sup>

Classification of the Rare Earth Elements.—In dealing with this subject it is convenient to commence by a discussion of how the elements of the rare earths fall into line with the most popular of the classifications of the elements, namely, the Periodic Classification as arranged by Mendel/eff

<sup>&</sup>lt;sup>1</sup> The most satisfactory determinations recorded in the pieceding table are undoubtedly the determinations of the atomic weights of needymium and prascodymium by Baxter and his co-workers. For more complete discussions of the atomic weights of the rare earth elements, the reader is referred to Clarke, A Recalculation of the Atomic Weights ("Smithsonian Miscellaneous Collections," vol 54, No. 3, 1910); Urbain, Bull Soc chim, 1909, [iv.], 5, 133; and the various articles by Brauner in Abegg, Handbuch der anorganischen Chemie, vol. 3, pt. 1, 1906.

Let it first be supposed, then, that the Periodic Table as usually drawn up 1 is correct. Then, since it must presumably be supposed that the eighth horizontal row contains three transition-elements in the Group VIII space, there are twenty vacant spaces between barrum and tantalum, waiting to be filled by one mert gas and nuncteen other elements - For these vacancies the atomic weight range is such that it includes the atomic weights of the rare earth elements, yttimin being excepted? The rare earth elements must therefore be placed in these vacancies, for, on the one hand, no other places are available, and, on the other, if other places were available, those in question would still have to be filled, and it is difficult to believe that nineteen metals of suitable atomic weights still await discovery

It requires, however, but a slight knowledge of the chemical and physical properties of the rare earth elements to realise that when these elements are placed in position in the table as indicated above, it can no longer be said that the properties of the elements are periodic functions of their atomic weights, i.e the Periodic Law breaks down. Such being the case, it seems hardly worth while to attempt any detailed discussion of the precise places the rare earth elements must occupy in the table, and therefore only a few words will be said upon the subject. The best arrangement appears to be the following -3

|     |    | I.  | 11  | 111 | IV | v    | vi   | VII | viii             |
|-----|----|-----|-----|-----|----|------|------|-----|------------------|
| VI  | 11 | Λд  | ea. | In  | Sn | Sb   | Te   | 1   | - · <del>-</del> |
| 11  | П  | Cs. | Br  | La  | Ce | Pr   | Nd   | -   |                  |
| -IX |    | Sm  | Eu  | Gd  | Ть | Dy   | - Ho | Er  |                  |
| X.  | •  | Tm  | Yb  | Lu  | Ct | Ta - | W    |     | Os, It, Pt       |

This arrangement is in harmony with Moseley's work on the high frequency spectra of the elements, according to which the sequence of atomic numbers is as follows

the blanks denoting missing elements 5. It also agrees with the observations of Hicks,6 (1) that the line spectra of Cd, Eu, and Hg exhibit triplet series of the sharp and diffuse type, the triplet intervals being approximately proportional to the squares of the atomic weights, (ii ) that Gd exhibits doublet series similar to

<sup>1</sup> See Frontispiece

<sup>2</sup> It has been already pointed out that yttrium may readily be found a place in the table;

that see a nearly pointer out that yetrium may readily be round a place in the table; see p. 238

Rydberg, Lunds Univ Arvskrift, N.F., Afd. 2, 1913, 9, No. 18, J. Chim. phus., 1914, 12, 585., Rev. gen. ser., 1914, 25, 734. Phil. May., 1914, [vi.], 28, 114. Hicks, Phil. May., 1914, [vi.], 28, 139. This arrangement differs considerably from that given by Brauner, Zeitsch. Erkhrichem., 1908, 14, 525. Rydbirg, however, prefers to repect Mendoléedl's arrangement and adopt an ingenious table in which the elements are arranged in four series, containing respectively 4, 15, 35, and 64 elements (1 $p^2$ , where p=1, 2, 3, and 4). His table is reproduced in Hicks's paper

<sup>&</sup>lt;sup>4</sup> Moseley, Phil. Mag, 1914, [v1], 27, 703, see also pp. 312, 363, and Vol. I.

Chap. VIII

It cole ments Moscley did not examine
Hicks, Phil. Trans., 1912, A, 212, 58.

Moscley also writes Ho in mistake for Dy.

those of Al, Ga, and Ti, and (iii.) that Sc, Y, La, and Yb 1 also show doublet series. It cannot, however, be reconciled with the other physical and the chemical properties of the elements, but this applies to every other similar arrangement. It should be pointed out that the arrangement given assumes that terbinin and thulium are true elements, points that have been disputed, and that Marignac's "ytterbum 'is composed of three elements, ytterbum, lutecum, and celtum.2 Moreover, it is necessary to assume that Series VIII. contains no transitionelements, and that an unknown rare earth element exists, the atomic weight of which hes between the values for needymmm and samarium.

So far it has been assumed that Mendeléeff's arrangement into groups and series is correct, and it is evident that the assumption is by no means satisfactory, since of the rare earth elements only yttrium, lanthanum, and ceruin can be readily fitted into the scheme. It is not surprising, therefore, that attempts have been made to improve matters by suitably modifying Mendeleeff's arrangement. Those attempts which alter that arrangement least are based on the assumption that all but one or two of the rare earth elements may be placed en bloc in a single place in the table 3 The arrangement proposed by Benedicks, for instance, is as follows --

| ,     |    |     |    |      |                                    |    |     |     |            |  |
|-------|----|-----|----|------|------------------------------------|----|-----|-----|------------|--|
| 1 1   | 0  | I   | П  | III. | IV.                                | v. | VI. | VII | VIII.      |  |
|       |    |     |    |      | <del></del>                        |    |     |     |            |  |
| VIII. | Xe | C's | Ba | Lι   | IV.<br>Ce, Pı, Nd, Sm, En, Gd, etc | Ta | W   | -   | Os, II, Pt |  |

and has the obvious disadvantage of including most of the rare earth elements in Group IV instead of in Group III as their valency indicates. hand, to group all the rare earth elements except cerrum in the Group III position is to depart from the atomic weight sequence - Brauner's 1902 proposal was expressed in terms of an hypothesis the precise meaning of which is not very clear, indeed, it has been said that it "belongs to the realm of puro fantasy". Stripped of its hypothetical garb, however, the proposal is identical with that advocated by Benedicks

At the time when they were published, the preceding proposals could only be viewed as makeshifts, similar to the arrangements of the transitionelements in sets of three in Group VIII - Essentially the same idea, viz. that the rare earth elements form a transition group, is expressed in the arrangements of Steele and Werner, although these chemists modify the nature of Mendeleeff's table considerably 5 Modern work on the radio-elements, however, has shown that it is often necessary to assume that several elements occupy one and the same place in the Periodic Table, an assumption analogous to the preceding However, theoretical views concerning the structure of the atom indicate that this only arises when several elements possess the same "atomic number," and the view that the rare earth elements have the same

<sup>&</sup>lt;sup>1</sup> Old ytterbunn (Yb, Lu, and Ct) (?).

<sup>2</sup> These questions are discussed later (pp. 358-365).

<sup>3</sup> Brauner, Chem. Zentr. 1900, 11-521; J. Russ. Phys. Chem. Soc., 1902, 34, 142;

Zeitsch annug. Chem., 1902, 32, 1; Bonedicks., ibid., 1904, 39, 41, H. Biltt, Ber., 1902,

35, 562; R. J. Mevel, Naturevsenschaften, 1914, 2, 781. See also Retgers, Zeitsch.

physical Chem., 1895, 16, 651.

<sup>4</sup> Wyrouboff and Venneul, Ann. Chem. Phys., 1906, [v1], 6, 166

<sup>5</sup> Steele, Chem. News., 1901, 84, 245; Weiner, Ber., 1905, 38, 914. New Ideas on Inorganic Chemistry, Irans. by Hedley (Longman & Co., 1911).

<sup>6</sup> See Vol. 1. Chap. VIII

"atomic number" 1 cannot be reconciled with Moseley's observations on high frequency spectra 2

So far, then, as a comprehensive classification, including all the known elements, is concerned, the rare earth metals offer formidable difficulties The problem of the interclassification of the rare earth elements is not in a much better position, judged from the purely scientific standpoint

The early division into cerite and gadolimite earths, based upon their mode of occurrence, dates from the time when but six of these earths were recognised. It divided the rare earth elements into two groups, the metals of the cerum and uttrium groups respectively. The subsequent discovery of the remarkable complexity of three of these earths was also associated with the observation that the properties which distinguished the cerite and gaelolimte earths failed to effect a sharp division into two groups, and an intermediate group of terbia earths was recognised, particularly by Marignac and Lecoq de Bo shandran - The importance of this group, however, as Urbain has remarked, was little appreciated for a considerable number of years, an error that contributed appreciably to the remarkable confusion into which the chemistry of the rare earths was thrown throughout the period 1878 to 1900,

With the recognition of the terbir curths as an intermediate group, it is possible to divide the rare earth elements into three groups, as follows.

| Cerium Group | Terbinin Group | Yttrinin Group |
|--------------|----------------|----------------|
| Lanthanum    | Europann       | - Dysprosium   |
| Cerum        | Gadohmum       | Holiounn       |
| Prascodymmm  | Teeb unc       | Yttmm          |
| Neodyminm    |                | Erbunn         |
| Samarum      |                | Thulum         |
|              |                | Ytterbunn      |
|              |                | Luteemm        |
|              |                | Celtum         |

The yttimm group is sometimes divided into parts, the yttimm group proper (Y, Yb, Lu, Ct) and the erbanic group (Dy, Ho, Tin, Er)

The following differences may be noticed between the properties of the compounds of these three groups

- (i) The platmocyanides of the cerum and terbinin groups are similar in composition, but differ in crystalline form, colour, and degree of hydration from those of the yttimm group (p. 272)
- (ii ) The double compounds MiCNS) 3Hg(CN), 12H<sub>2</sub>O are monoclinic in
- the comm, and trichme in the vitrum group (p. 272) (m.) The subcotungstates of the formula  $M_4(W_{12}SiO_{10})_q.81H_2O$  and the acid subcotungstates of the formula  $MH(W_1.SiO_{10}).18H_2O$  are missing in the yttimin group, while the acid salts of the formula 2MH(W12SiO40) 49H2O are missing in the cerum group (p. 266)
- (iv) The intrates of the metals of the cerum and terbium groups form double intrates with the intrates of inn- and bi valent metals, but the nitrates of the metals of the yttrium group do not (p. 269)

Other double compounds of the nitrates may be also divided into groups

<sup>&</sup>lt;sup>1</sup> Suggested by van den Brock, Phil. Mag., 1914 [vi.], 27, 455.

<sup>2</sup> The position of the rate earth elements in the Periodic Table is also discussed by Rudof (Ealsek, anong, Chem., 1903, 37, 177), and by Harkins and Hall (J. Amer. Chem. Soc., 1916, 38, 169).

<sup>3</sup> Urbain, Man. Chim. Phys., 1900, [vii.], 19, 184

Thus the compounds with antipyrine appear to be of the type M(NO<sub>3</sub>)<sub>8</sub>.

 $3C_{11}H_{12}ON_2$  in the cerum and  $M(NO_3)_3$   $4C_{11}H_{12}ON_2$  in the yttrium group. (v) The double sulphates of potassium and the metals of the cerum group are insoluble in a cold, saturated solution of potassium sulphate, in the case of the terbium group they are sparingly soluble, and in the case of the yttrium group readily soluble (p. 261)
(vi.) The double carbonates of the alkali metals and the metals of the

cerium group are decomposed by water and dilute alkali carbonates, but not by concentrated alkali carbonates, the corresponding double salts in the yttrum group are readily soluble in and not decomposed by dilute alkali carbonates (p. 271)

(vii) The oxalates of the metals of the cerium group are only slightly soluble in hot, concentrated ammonium oxalate, those of the vttrium group are more readily soluble, and double salts may be crystallised from the solutions (p. 271).

The preceding differences are not of a very fundamental character Moreover, differences of a semilar character may be pointed out which are not in accordance with the classification into the cerium, terbium, and

In other natural families of elements, e.g. the alkali or the alkaline earth metals, progressive variation in properties usually follows the order of increasing atomic weights. The same statement cannot be so confidently affirmed in the case of the rare earth elements, for yttrium is then completely removed from the elements which chemically it most closely resembles, namely, holmium and erbium - Excluding yttrium, however, it may be stated that the solubilities of corresponding salts usually follow the order of the atomic weights of the rare earth elements (see p 325), that the atomic weight sequence places the terbium group between the cerium and ottrium groups, and that the same arrangement brings samarium and europium, the only two rare carth elements known to form derivatives of the type MA, next to one another. There are, however, several indications that the atomic weight sequence is often associated not with a progressive, but with a periodic variation in properties, see, for instance, the magnetic susceptibilities of the oxides (p. 257), the melting-points of the chlorides (p. 253), and the molecular volumes of the ethylsulphates (p. 279) 3

# COMPOUNDS OF THE RARE EARTH ELEMENTS.

The great majority of these compounds are salts derived from a series of basic sesqui-oxides, M2O, Three lower halides are known, viz. SmCl., SmI2, and EuCl, in which the metals may be bivalent, but the molecular weights of the compounds are not known. Cerum forms a series of compounds in which it is quadrivalent, presumably praseodymium is also quadrivalent in the dioxide PiO2.

Kolb, Zeitsch anorg Chem , 1913, 83, 113

See, e.g., the compositions of the schaeates and 1 · 4 : 2-biomonitiobenzenesulphonates

<sup>(</sup>p. 281).

See also Jacger, Proc. K. Akad. Wetensch. Amstradam, 1914, 16, 1095, Rec transchim, 1914, 33, 342; R. J. Meyer, Naturoussenschatten, 1914, 2, 781. A few physical properties appear to be anomalous, eq. the molecular volume of cerous ethylsulphate (p. 279) and the molecular volumes of the double intrates of neodymium and prescodymium of the type  $3M''(NO_3)_2 2M'''(NO_3)_2 21H_2O$  (p. 270).

So far as the derivatives of the sesqui-oxides are concerned, the rare earth elements appear in their chemical relations to be intermediate between the alkaline carth metals and magnesium, while also bearing some resemblance to iron, alimmium, and chromium The colours of the salts are as follows .-

Lanthanum, colonrless. Cerrum (ous), colourless Prascodymum, green. Neodymum, red to reddishviolet Samarum, topaz yellow.

7. 1

Europium, pale rose. Gadolimum, colourless. Terbium, colomiless. Dysprosium, yellow to greenish yellow Holmum, yellow to orange.

Yttrum, colourless. Erbmm, deep rose Thulmm, green. Ytterbuum, colom less. Lutecum, colom less.

In a general way it may be said that the salts have the properties that would be anticipated of salts derived from fauly strong bases. Of the commoner salts, the fluorides, iodates, chromates, phosphates, carbonates, and oxalates are practically insoluble in water at the ordinary temperature, and the sulphates are but sparingly soluble. Corresponding salts of the rare earth elements are frequently isomorphous; numerous instances are cited in the succeeding page

It is often found that when corresponding salts of the rare earth elements are arranged in order of solubility, the order is identical with the order of atomic weights of the contained metals, with the exception of yttinum, which comes between holimum and eibium. This matter is discussed more fully in the next chapter (see p. 325). The order of arrangement La, Ce, Pr. Nd, Sm, Eu, Gd, Th, Dy, Ho, Y, Er, Tm, Yb, Lu, Ct is sometimes conveniently spoken of as the serval order of the care earth elements. With reference to the solubilities of the rare earth salts, it is interesting to note that several series of salts exhibit the uncommon property of being less soluble in hot than in cold water, e.g. the hydrated sulph ites, double sulphates, and dimethylphosphates.

With regard to the formation of double salts, the rare cuth compounds present several points of interest. Thus, numerous double initiates are known, although outside the field of rare cuth chemistry such salts are very uncommon. Of more interest are the double salts containing two different acid radicles, eg lanthanum ovalochloride,  $\operatorname{La_2(C_2O_4)}$ ,  $\operatorname{LaCl_5} 7\frac{1}{2}\operatorname{H_2O_5}$ or 2La(C,O) Cl 5H,O as it is more conveniently written. In addition to ovalochloudes, there are known ovalountrates, ovaloboundes, ovalo-bodides, ovalosulphates, intratosulphates,  $M(NO_2)\,SO_4\,H_2O_5$ , and chloroeacodylates,  $2M[(CH_3)_2\,AsO_2],M(H_4\,18\,H_2O)$ 

# THE RARE EARTH ELEMENTS AND HADROGEN

Hydrides, MH, -The rare earth metals combine directly with hydrogen. as Winkler and Matignor have shown? The latter observed that the hydrides

<sup>&</sup>lt;sup>1</sup> For measurements of the conductivities of the carth salts, see Muthmann, Rev., 1898, 31, 1829, Ley, Zeits h. physikal. t. hem., 1899, 30, 193, Rimbach and Schubert, ibid., 1909, 67, 183, Aufrecht, Inaugural Description (Berlin, 1904). Jones and Reese, Amer. Chem. J., 31, 1528, key, zeroson program Description (Berlin, 1904). Jones and Reese, zimer vierne, s., 1898, 20, 606, Jones and Allen, ibid 1896, 18, 321, A. A. Noyes and J. Johnston, J. Amer. Chem. Sec., 1909, 31, 987, Holmberg, Inkins Kem. Min. God, 1903, x, 1, Jantsch and Grunkiaut, Zeitsch amorg. Chem., 1913, 79, 305, A. Heydweiller, Zeitsch physikal. Chem., 1915, 89, 281. For a selection of numerical results, see under the headings of the

Various metals.

<sup>2</sup> Winkler, Ber, 1891, 24, 873, 1966 (La, Ce, Y), Matignon, Compt. rend., 1900, 131, 891 (La, Ce, Nd, Pr, Sm).

are readily dissociated by heat. Muthmann, Kraft, and Beck have prepared various hydrides by the direct union of metal and hydrogen at 220°-270°, and find that these compounds are of the type MIL, except the neodymnum compound, which is NdH21. The hydrides are brittle, amorphous solids, the cermin and lanthanum compounds being bluish-black, the neodymnum compound indigo blue and the praseodymium compound green in colour atomic heat of hydrogen in ceruum and prascodymium hydrides is 2 1.2

#### THE RARE EARTH ELEMENTS AND THE FLUORINE GROUP.

Fluorides, MF, -The anhydrous fluorides are produced by the action of fluorine on the carbides (Moissan), and by heating the oxides to redness in a current of hydrogen fluoride. Hydrated fluorides may be prepared by double decomposition. According to Hirsch, the best procedure consists in adding concentrated hydrofluoric acid solution to a hot solution of the chloride containing very little free hydrochloric acid. The precipit ded fluoride is washed by decantation, twice with hot water and several times with 95 per cent alcohol, and centrifuged till practically free from alcohol. The anhydrous fluoride may then be obtained by covering the product with absolute alcohol, evaporating to dryness, and heating the residue first at 100°, and finally at 200°.

The precipitated fluorides are practically insoluble in water and dilute mineral acids. In a large excess of hot, concentrated hydrochloric acid they are perceptibly soluble, the fluorides of the cerum group being more soluble than those of the yttrium group. In similar circumstances the fluorides of thorum and scandium are much less soluble?

The fluorides of the rate earth elements crystallise in the cubic system and are isomorphous with fluorspar, in which mineral they frequently occur in traces (see pp. 217, 234)

Chlorides, MCl, -- The anhydrous chlorides may be prepared in the pure state by the following methods

- (1) Dehydration of the hydrated salt, by heating the latter to 105 -130° in a current of pure, dry hydrogen chloride (taking care to avoid fusion of the hydrated salt) until no more water is given off, and then heating the residual monohydrate, MCL, II,O, to 180-200' in the stream of gas as before, When water ceases to be given off, the temperature may be raised until the chloride fuses 6
- (ii.) Conversion of the oxide (sulphate or oxalate) into chloride by heating it in a current of disulphur dichloride vapour? The process is started at 250° and the temperature slowly raised to a point about 50 below the . . . . . . . . . . . . .

- --

Muthmann and Kraft, Innalen, 1902, 325, 261 (La, Ce), Muthmann and Beck, ibid, 1904, 331, 58 (Nd, Pr), see also Muthmann and Baur, Innalen, 1902, 325, 281 (La, Ce).
 kellenberger and Kraft, Innalen, 1902, 325, 279. Temperature runge not stated.
 Van Hasgen and E. F. Smith, J. Inner Cham Soc., 1911, 33, 1504 (La, Ce, Y).
 Hirsch, J. Ind. Eng. Chem., 1911, 3, 885.
 See under Scandum, p. 207.
 Mattgnon, Jan. Chim. Phus., 1906, [vin.], 8, 364 (general), 243 (Nd), 386 (Pr), 402 (Sm), 426 (La), 433 (Y), 410 ("Yh"). Compt. rend., 1901, 133, 289, 1905, 140, 1637 (Nd); 1902, 134, 427 (Pr), 1908 (Sm. Y, "Yh"). Baxter and Chapin, J. Amer. Chem. Soc., 1911, 33, 104, Baxter and Stewart, bid., 1915, 7, 516 (Pr), cf. Mariginge, Ann. Chim. Phys., 1853, [in.], 38, 148, Hermann, J. prokt. Chem., 1861, 82, 385.
 Not a maxime of sulphur chloride and chlorine, since in that case the product always contains oxy, bloride that cannot be converted into chloride.

contains oxychloride that cannot be converted into chloride.

melting-point of the chloride. Finally, the chloride is fused and coded in a current of dry hydrogen chloride 1. A rapid method of preparation, which is essentially a combination of methods (i) and (ii), consists in evaporating a solution of the chloride to divness on the sand-bath and heating the solid residue in a current of dry hydrogen chloride and disulphur dichloride

(in ) Conversion of the oxide into chloride by heating it in a stream of carbonyl chloride2 or a mixture of carbon monoyide and chlorine,3 or in a stream of carbon tetrachloride vapour 1

(iv) Conversion of the oxidate into chloride by heating it carefully in a stream of dry hydrogen chloride? The oxide may be similarly converted into the chloride at a red heat, but it is practically impossible to carry out the conversion quantitatively 6

The anhydrous chlorides are formed when the metals are heated in chlorine or hydrogen chloride, when the callides are heated in chlorine, and when the sulphides are heated in chlorine or hydrogen chloride, but these processes are of little value as methods of preparation, except, perhaps, the sulphide method. Further, the classic method of heating a mixture of the oxide and carbon in a stream of chlorine is of no practical value, since the chlorides are non volatile at a red heat

The anhydrous chlorides are solids which melt at a red heat and crystalliso The densities (at 0) and melting points are given by Bourion as on cooling follows (melting-point of yttirum chloride by Matignon) -

```
LaCl_{i} = CcCl_{i} - PrCl_{i} - \nabla dCl_{i} - SmCl_{i} - GdCl_{i} - TbCl_{i} - DvCl_{i} - Vcl_{i}
                                111 127 152
Density
            379 392 107
  4 35 3 67
            890 848 810
                                781 686 628
  588 - 680 - 686
Mpt C.
                                 YEC 10 Luci "
                     Density (
                                  3.98
  , 916
                                  850
```

According to Matignon, the densities at 18 are as follows

The volatility of the oblorides at a rol heat is very slight, but the oblorides of the yttrium group are distinctly more volatile than those of the cerium group

<sup>&</sup>lt;sup>1</sup> Bourion, Inn. Chim. Phys., 1919, [viii], 20, 547, 21, 19 Ta. Ce. Fr. Nd., 84, Gd., Tb., Dv., Yb., Lii., & Matignen and Boarion, Compt. cond., 1904, 138, 641. Matignen, that, 1905, 140, 1181, 1839.

"Chairent, Compt. rend., 1911, 152, 87.
'Didien, Compt. rend., 1885, 101, 882., Ann. Physic. Normath., 1887, [iii] 4, 65; Duboin, that, 1888, [iii], 5, 416., Matignen and Delepine, Compt. rend., 1911, 132, 37.
'Demaryay, Compt. rend., 1887, 104, 111. Quantin, that, 1887, 104, 22), 1888, 106, 1074; L. Meyer, Rev., 1887, 681, Matignen and Delepine, Le. of Robinson, Proc. Roy. Soc., 1884, 37, 150., Chem. Nais, 1884, 50, 251 (Ce).
Matignen, Ann. Chem. Phys., 1906, [viii], 8, 361. Petterson, Anker. annu Chem., 1893, 41.

<sup>1893, 4, 1.

7</sup> Hillebrand and Norton, *ibid.*, 1875, 155, 633

8 Petterson, *liet.*, 1895, 28 2419, Morssan, *compt. read.*, 1896, 122, 357, 1900, 131, 595, 924. Morssan and Étard, *ibid.*, 1896, 122, 573

9 Mothmann and Statzel, *Ber.*, 1899, 32, 3413

10 Metaral of doubtful marity.

<sup>10</sup> Material of doubtful purity.

The chlorides are extremely hygroscopic, and dissolve in water with the evolution of considerable heat The following values for the heats of solution are given by Matignon -

The aqueous solutions are neutral to methylorange. readily soluble in alcohol, less so in pyridine, and insoluble in other, chloroform, acetone, and the chlorides of phosphorus, arsenic, sulphur, and tin (Matignon); 100 grains of pyridine at 15° dissolve the following amounts of the anhydrous salts

The chlorides are readily prepared in solution by dissolving the oxides, hydroxides, or carbonates in concentrated hydrochloric acid, by concentrating the solutions over sulphuric acid, cooling the hot, saturated solutions, or saturating the cold aqueous solutions with gaseous hydrogen chloride, crystalline, hydrated chlorides are readily prepared, usually of the type MCl<sub>3</sub> 6H<sub>2</sub>O. The hydrates thus obtained, when heated to 105–130 in a stream of dry hydrogen chloride, are partly dehydrated, the monohydrates, MCl, II.O, being produced, the monohydrates are completely dehydrated at 180°-200° in hydrogen chloride 2. The hydrated salts cannot be dehydrated by simple heating in air without decomposition taking place

Caesium chloride combines with the rare earth chlorides, and the following crystalline salts have been obtained .— 3

Other double chlorides containing rare earth metals and the metals of Groups I. and II. are not known, with the exception of the aurichlorides. Double chlorides with antimony, bismuth, and tin chlorides, and the chlorides of platinum, have, however, been prepared. The chloroplatinates are of interest, since their composition affolds evidence for the terralency of the rare earth elements (see p. 237). They are readily soluble, orange-red, hygroscopic, crystalline salts, and the amount of water of crystallisation they contain is variously given as 12, 125, and 13H,O. The chloroplatmates of the metals of the cerum group are isomorphous, crystallising in tetragonal bipyramids - 6

$$\begin{array}{lll} {\rm CeCl_3.PtCl_4.12H_2O} & a.c = 1 & 1.1272 \\ {\rm PrCl_3.PtCl_4.12H_2O} & = 1 & 1.13 \end{array}$$

Bainobey, J. Amer. Chem. Soc., 1912, 34, 1174.
 Matignon, see it ferences on p. 252.
 R. J. Meyer and Wassjuchnow, Zeitsch. anorg. Chem., 1914, 86, 257.
 Dehnicke, Inaugural Dissolution (Beilin, 1904).
 See Vol. IX., or Nilson, Bull. Soc. chim., 1877, [ii.], 27, 206, 208; also Nilson, ibid., 1877, [ii.], 27, 210, for the chloroplatinites
 Marignao, Ann. Chim. Phys., 1873, [iv.], 30, 65; Topsoe, Bihang. K. Swinska Vet.-Akad. Handl., 1873-5, 2, No. 5, Sodeistrom, Zeitsch. Kryst. Min., 1902, 36, 194 (Pr).

The rare earth chlorides also form double compounds with mercuric cyanide, of the type RCl,  $3\mathrm{Hg}(\mathrm{CN})$ ,  $8\mathrm{H}_2\mathrm{O}$ , these compounds lose  $8\mathrm{H}_2\mathrm{O}$  over concentrated sulphune acid or at 100° 1

The chlorides of the rare earth elements form compounds with alcohol, pyridine, and other organic bases. The compounds tabulated below have been obtained by Matignon —2

| PiCl, 2C,H,N                                       | NdCl, 3C,H, OI                       |
|----------------------------------------------------|--------------------------------------|
| NdCl <sub>1</sub> 3C <sub>2</sub> H <sub>2</sub> N | NdCl, C, Ĥ, ŎH                       |
| SmCl, 3C,H,N                                       | YCl <sub>a</sub> Č,Ñ <sub>a</sub> ÓH |
| YCl <sub>1</sub> 3C <sub>2</sub> H <sub>5</sub> N  | 3 . 3                                |

Oxychlorides, MOCI —These compounds may be prepared, sometimes in the crystalline form, by passing a slow current of air or steam (carried by an mert gas) over the molten, anhydrous chlorides 3. They are also produced by heating the hydrated chlorides, or better, the ovalochlorides,4 M(C2O1)XI dI2O, to a red heat The best method for obtaining them in the crystalline state consists in adding the oxides or sulphates to molten magnesium chloride, allowing to cool, and extracting the soluble salts with dilute acetic or hydrochloric acid 5

The oxychlorides are practically insoluble in cold water, dissolve very slowly in dilute acids, and are converted into oxides by prolonged heating in an They are of considerable practical value, since their reflection spectra (p 291) are very well defined. For the purpose of examining these spectra the oxychlorides are readily prepared in the following manner. The oxides are dissolved in hydrochloric acid, ammonium chloride added, the solution evaporated to diviness, and the residue heated carefully 6. It is then extracted with water and the insoluble spangles of oxychloride collected on a filter paper 7

Bromides, MBi .- These salts have not been examined so closely as the chlorides. The anhydrous bromides have been prepared by heating the anhydrous chlorides in pure, dry hydrogen bromide, taking care to avoid fusing during the operation, by heating the sulphides in dry hydrogen brounde, by heating the oxides in sulphur bromide vapour, 10 and by heating the oxides in a current of sulphur chloride and hydrogen brounde, the latter being in excess 11. The bromides resemble the chlorides in properties. They are shightly soluble in acetone 12. From aqueous solution they usually crystallise as hexalydrates MBr, 6H2O

Iodides, MI3 -Very little is known of these salts - Matignon has prepared the anhydrous rodides of neodymnum, prascodymnum, and samarrum by

Alén, Bull Soc chim, 1877, [n], 27, 305 (La. Ce, "Df," Y, "E,")
 Matignon, references on p. 252, see also R. J. Meyer and Koss, Ber., 1902, 35, 2622;
 Jefteson, J. Amer. Chem. Soc, 1902, 24, 540, Hartwell, chid., 1903, 25, 1128.
 Barbieri and Calcolari, Atti. R. Accad. Leavis, 1911, [v], 20, 1, 164.
 Matignon, references on p. 252 (Pr, Nd, Sm), Bourton, Ann. Chim. Phys., 1910, [vnn.], 27, 40 (Zd. Liv.)

<sup>21, 19 (</sup>Gd, Dy)

4 Job, Compt. rend., 1898, 126, 246.

5 K. A. Hofmann and Howhele, Ber., 1914, 47, 258 (Pr. Nd, Sm, Er).

6 K. A. Hofmann and Howhele, Ber., 1914, 47, 258 (Pr. Nd, Sm, Er).

b It is advisable to add a mixture of sodium and potassium chlorides to the original solution and heat finally until the mixed chlorides fuse.

7 Usbarra for Charles 1999 Lead to the corrections of the charles fuse.

Urbain, Ann Chim. Phys., 1900, [vii], 19, 222.
 Matignon, references on p. 252 for Nd and Pr.
 Muthmann and Stutzel, Ber., 1899, 32, 3413 (Ce).
 Barre, Bull. Soc. chim., 1912, [vi], 11, 433.
 Bourton, Compt. rend., 1907, 145, 243 (La, Ce, Nd, Pr, Sm, Gd, Dy, Tb, "Yb")
 Barnebey, J. Amer. Chem. Soc., 1912, 34, 1174.

heating the anhydrous chlorides in pure, dry hydrogen iodide and avoiding fusion during the process. They are extremely hygroscopic, and soluble in acctone 2

Bromates, M(BiO<sub>3</sub>), ... The biomates are best prepared by covering barrum bromate with water, heating on the steam-bath, and adding gradually, with continual stiring, the neutral raic earth sulphates. A slight excess of the barum bromate is necessary. When the double decomposition is complete, the barium sulphate is removed by filtration and the biomates crystallised from the solution They form an isomorphous series of salts of the general formula M(BrO<sub>D)</sub>,911,0, crystallise in hexagonal prisms, and are easily soluble in water. The melting-points and solubilities at 25 (in grams of crystallino salt dissolved by 100 grams of water) of a number of bromates are as follows : - 1

Iodates, M(10), - The rodates are readily prepared by double decomposition They are crystalline salts, sparingly soluble in water, but readily soluble in concentrated intricated. In the latter respect they differ markedly from the iodates of zincommit and thornum  $^{5}$ 

#### THE RARE EARTH ELEMENTS AND THE OXYGEN GROUP.

Sesqui-oxides, M.O. - Each of the rare earth elements forms a basic sesqui-oxide, the series of sesqui-oxides constituting the rare earths themselves With the exception of the compounds of cerum, praseodymum, and terbinin, the sesqui-oxides are stable in an even at a bright red heat, and so may be produced by the ignition of the hydroxides, carbonates, intrates, ovalates, sulphates, and many other salts of oxyacids. The sesqui-oxides of prascodymium and terbuun, however, combine with oxygen when heated, forming dark brown peroxides of somewhat indefinite composition, these peroxides may be reduced to sesqui-oxides in a stream of hydrogen at a red heat. Cerum sesqui oxide, on the other hand, is extremely easily oxidised to cermin dioxide, which can only be reduced again with great difficulty

When prepared by the ignition of smitable salts, the rare earths are obtained as amorphous powders; lanthana has also been obtained in the crystalline form. The appearance and chemical reactivity of a rare earth depend upon its method of formation, i.e. whether it has been formed from the oxalate, say, at a red heat or from the sulphate at a white heat. Practically nothing is known as to the cause of this difference, it does not even appear to be known whether the different forms have different densities As a rule, the differences in reactivity are attributed to the fact that the

various modifications are highly, but unequally, polymerised.

The seson oxides are only reduced to the metallic state with considerable

Matignon, teferences on p. 252
 Barnebey, J. Amer. Chem. Soc., 1912. 34, 1174.
 Rammelsberg, Poop. Annaten, 1812, 55, 61, Marignac, Ann. Min., 1859. [v.] 15, 221.
 James, J. Amer. Chem. Soc., 1908, 30, 182; Chem. News, 1909, 97, 61, James and Langelier, J. Amer. Chem. Soc., 1909, 31, 913, Chem. News, 1909, 100, 85
 Davis, J. Amer. Chem. Soc., 1889, 11, 26, R. J. Meyer and Spetci, Chem. Zeit., 1910, 34, 306; R. J. Meyer, Zeitsch. anorg. Chem., 1911, 71, 65.

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difficulty, since their heats of formation are very high. The following values for the heats of formation (in Cals per gram-equivalent of oxide) of lanthana, praseodymia, neodymia, ceria, and the refractory oxides magnesia and alumina, will serve to illustrate this fact -1

The magnetic susceptibilities of the rare earths are of considerable practical importance, since the values vary very considerably and thus afford a rehable means of following the processes of fractional crystallisation, precipitation, otc, (see p 330) The following values have been recorded for the magnetic susceptibilities (x) per grain of sesqui-oxide at the ordinary temperature —  $\frac{1}{2}$ 

The values for lutecia and celtia are only approximately known, ytterbia being about four or five times as paramagnetic as luteria, which in its turn is four times as paramagnetic as celtia. The values for dyspresia, holima, erbia, thulia, and ytterbia are known to be in descending order of magintude. Prascodymia is not so paramagnetic as neodymia. It is of interest to note that the three elements which may definitely be assigned positions in Subgroup III. A of Mendeleeff's Table, viz scandium, yttrium, and lanthanium, all form feebly diamagnetic oxides

The rare earths are fauly strong basic oxides, being much stronger bases than the sesqui-oxides of Subgroup III B From general chemical and thermochemical i considerations it may be concluded that they are stronger bases than magnesia, but weaker than the alkaline earths. Thus, the strongly ignited oxides are readily soluble in mineral acids,4 several of them combine directly with water and absorb carbon dioxide from the air, and all of them liberate ammonia from cold solutions of ammonium salts. Normal salts of numerous weak acids, e.g. chromic, carbonic, and acetic acids, are readily prepared. The salts derived from strong acids are only hydrolysed to a very slight extent even in dilute aqueous solution at the ordinary temperature.5

The determination of the relative strengths of the rare earths as bases has so far only been attempted in a qualitative manner, and the results are

<sup>&</sup>lt;sup>1</sup> Muthmann and Weiss, Annalen, 1901, 331, 1
<sup>2</sup> Urbain and Jantsch, Compt. rend., 1908, 147, 1286 (Nd. Sm., Eu, Gd., Th., Dy). Urbain, ibid., 1909, 149, 37 (Tb). Blumenteld and Urbain, ibid., 1914, 159, 323 (Yb). R. J. Meyer and Wuonnen, Zeitsch. annary. Chem., 1913, 80, 7 (La, Y). See also S. Meyer, Sitzmuyaber, K. Akad. Wiss. Wiss., 1900, 10, 11a, 492, 1902, 111, 11a, 38, Aomatsh., 1892, 20, 369; 1908, 29, 1017. Ber., 1900, 33, 320, 1918. Ann. Physik, 1900, [iv.], 1, 664, 668., Onnes and Perrier, Proc. K. Akad. Wietensch. Amsterdam, 1912, 14, 115., Onnes and Oosterhus, abid., 1912, 15, 322. On the magnetic dichroism of the rare cattles, see Meslin, Compt. rend., 1909, 148, 1598.
<sup>2</sup> Mathgroon, Ann. Chim. Phys., 1906, [vin.], 8, 212, 364, 386, 402, 426, 433, 440; 1907, [vin.], 10, 104.

Mangnon, Ann them Thys., 1808, 1809, 1809, 1909, 1909, 1909, 100, 104

The most basic oxides are more readily soluble than the others

See Muthmann, Ret., 1808, 31, 1829, Lev. Zetsch physikal Chem., 1899, 30, 193;
Rimbach and Schubert, ibid., 1909, 67, 183, Denham, Zeitsch. anorg. Chem., 1908, 57, 378, Autrecht, Inaugural Dissertation (Berlin, 1904).

somewhat contradictory. The strongest base is lauthana, the earths of the cerium group are stronger than those of the yttimm group, yttma itself being excepted, and the terbia earths are intermediate in strength between the cena and yttria earths. Yttria itself is possibly as strong a base as neodymia. The rare earth elements are most probably arranged in the order of decreasing basic strength of their oxides as follows:-

This order is usually in harmony with (i.) the order in which the initiates decompose when heated, (ii ) the order in which the hydroxides are precipitated from the mixed salts by the gradual addition of animonium, sodium, or potassium hydroxide, and (iii) the relative degrees of hydrolysis of aqueous solutions of the sulphates. It is in complete disagreement with the order obtained by Wohler and Grunzweig 2 from measurements of the dissociation tensions of the anhydrous sulphates, it is difficult to believe that then results are accurate. Thermochemical data indicate that lanthana, prascodymia, neodymia, and samana are arranged in order of diminishing basic strength in the order in which they have been named. Thus, in the reaction --

$$[M_2O_8] + 6(HCI) = 2[MCI_3] + 3[11,O] + q Cals.$$

the values of q are -8

and in the reaction -

$$[M_2O_3] + 3[H_2SO_4] = [M_2(SO_4)_3] + 3[H_2O] + i$$
 Cals.

the values of r are -

In connection with the order of basic strength given above, there is one curious anomaly. Fractional precipitation with ammonia indicates that samaria is a stronger base than europia, but a weaker base than gadolima,5 while other methods indicate that samaria is stronger than gadolinia. It may be pointed out, however, that it is only an assumption to regard the order in which the hydroxides are precipitated as the order of their relative strengths, the weakest precipitating first. In accordance with the ionic theory, the order of precipitation represents merely the order of increasing solubility-products, and this only represents the order of mcreasing basic strength of the hydroxides provided that the hydroxides are very largely dissociated in solution and the un-ionised portions have approximately equal solubilities.

<sup>&</sup>lt;sup>1</sup> Katz and James, J. Amer. Chem. Soc., 1914, 36, 779.

<sup>8</sup> Wohler and Grunzweig, Ber., 1913, 46, 1726

<sup>\*</sup> Matignon, Loc et al. (1906, 142, 276

\* Lecoq de Borsbaudian, Compt. rend., 1890, 111, 393, 1892, 114, 575, Benedicks, Zeitsch. anorg. Chem., 1900, 22, 393.

Higher Oxides. - Cerrum forms a well-defined basic dioxide, CeO<sub>2</sub>; pracedymnum also forms a dioxide, PrO<sub>20</sub> and terhium forms a higher oxide of the approximate composition Tb<sub>4</sub>O<sub>7</sub>. These substances are not peroxides in the strict sense of the term, i.e. they are not constituted like hydrogen perovide. They are described under the headings of the respective metals

All the rare earth elements appear to be capable of forming true hydrated perovides of the type M2O5 aH,O, the constitutions of the compounds being expressed by the formula (OII), MOOII They may be obtained by the addition of hydrogen peroxide and ammonia to solutions of rare earth salts, and are described under the headings of the respective metals

Hydroxides, M(OH), - The hydroxides of the rare earth elements may be obtained as gelatinous precipitates by the addition of excess of ammonium, or an alkali hydroxide, (or ammonium sulplude) to hot, diluto solutions of the intrates, chlorides, etc. The hydroxides are insoluble in excess of precipitant, when a deficit of the latter is added, or the precipitation carried out with cold solutions, the precipitated hydroxides are contaminated with basic salts

The hydroxides are fairly strong bases, and absorb earbon dioxide from the an They interact readily with acids to produce salts

In the presence of ammonium acetate the hydroxides are only slowly and incompletely precipitated by the addition of ammonia, this property may be conveniently utilised in separating moderate amounts of non-from the rare earth elements  $^{1-}$ . The precipitation of the hydroxides may also be inhibited by the addition of citric acid and other organic hydroxy acids. The case of tartaric acid is of particular interest. When sufficient of this acid has been previously added, precipitation of the hydroxides by sodium hydroxide is completely inhibited. Potassium hydroxide, on the other hand, gives precipitates of double potassium tartiates with the members of the yttrium group, but only from boiling solutions. Lastly, ammonium hydroxide gives rise to precipitates of double ammonium tartiates with the yttrium group oven in cold solutions, the precipitates become crystalline when warmed? Muller 3 has found the precipitation of cerous and neodymnum hydroxides to be inhibited by the addition of glycerol

Sulphides, M.S. - The sulphides of the rare cutth metals cannot be prepared by wet methods. The best methods of preparation consist in heating either (a) the oxide or (b) the anhydrons sulphate to redness in a current of pure, dry hydrogen sulphide 1

The sulphides have been obtained as coloured powders, sometimes in the crystalline state They are comparatively stable in the an, but are slowly

<sup>&</sup>lt;sup>1</sup> Lecoq de Boisbaudian, Compt. cond., 1899, 111, 394, Sterba, Ann. Chem. Phys., 1904, [viii ], 2, 199, of the work of Delaloutain and others on colloidal basic arctates (Delafontaine, Chem. News, 1890, 73, 284, W. Bilte, Ber., 1901, 37, 719, Orboll, Chem. Zett.,

fontaine, Chem. Noves, 1890, 73, 284, W. Bille, Ber., 1901, 37, 110, Orion, Cason., 1907, 31, 45)

2 R. J. Meyer and Goldenberg, Chem. News, 1912, 106, 13, Meyer and Hauser, Dia Analys der selten n Erden und der Erdeauren Stuttgatt, 1912), p. 43.

3 Muller, Zeitsch anorg. Chem., 1905, 43, 320.

4 (a) Didier, Compt. rend., 1885, 100, 1461, 101, 882 (La, Ct.), Sterba, Jun. Chem. Phys., 1904, [vm.], 2, 193 (Ce.), (b) Muthmann and Statted, Ber., 1892, 32, 3413 (La, Ce.), Pr., Nd.; Sterba, loc. cit. (Ce.). Edmann and Witth, Annalen, 1908, 361–190 (Sm., Gd.) The carly Interature on the sulphides is as follows: Mosander, Popp. Annalen, 1820, 11, 406 (Ce.), Berimger, Annalen, 1842, 42, 138 (La), Lange. I. pakt. Chem., 1861, 82, 129 (Ce., La), Heimann, J. piakt. Chem., 1861, 82, 385 (La), Frenchs and Smith, Annalen, 1878, 191, 331 (La, Ce.), Popp, stud., 1864, 131, 197 (Y).

hydrolysed by boiling water to the hydroxides, hydrogen sulphide being evolved. The sulphides are readily decomposed by dilute mineral acids.

Oxysulphides and polysulphides are known only in a few instances Sulphites, M2(SO3)3 -The sulphites of the rare earth metals are practically insoluble in water, and may be obtained by double decomposition between sodium sulphite and rare earth salts 1 They are soluble in aqueous sulphurous acid, since the hydroxides, oxides, and carbonates may be dissolved in an excess of that reagent. The solutions thus obtained deposit crystals of the normal sulphites when evaporated over sulphuric acid, or when heated 2

Thiosulphates, M2(S2O3)3 -The rare earth elements form well-defined crystalline thiosulphates, which are readily soluble in water hydrolysed with the precipitation of basic salts when their aqueous solutions are boiled,3 in which respect they differ sharply from the corresponding salts of scandium, zirconium, and thorium, which may be completely hydrolysed.

Sulphates, M2(SO4)3 -The anhydrous sulphates of the rare carth metals may be prepared by heating the oxides, hydroxides, carbonates, chlorides, nitrates, or oxalities with an excess of concentrated sulphuric acid, and eliminating the excess of acid and destroying acid sulphates by licating to 500°. It is a matter of great difficulty to obtain the pure, anhydrous, normal sulphates by this method (see p. 240) An alternative method of preparing the anhydrous sulphates consists in dehydrating the hydrated salts at about 400°. At temperatures higher than 500°, decomposition commences, and at a white heat the sulphates are completely decomposed, the oxides being left.

The anhydrous sulphates of the rare earth metals are hygroscopic solids which dissolve in water with the evolution of heat - In order to obtain them in concentrated solution it is necessary to add the powdered solid gradually, with stirring, to ice-cold water, avoiding any local rise in temperature Con centrated solutions thus obtained are metastable with respect to the hydrated sulphates. Moreover, the solubilities of the hydrated salts diminish with rise of temperature very markedly 4 Hence, when a cold, concentrated solution of a rare earth sulphate is warmed, a copious precipitate of a hydrated salt is obtained, which cannot be brought into solution by cooling unless a considerable excess of the solvent be added. The most important hydrates of the sulphates are the octahydrates, M2(SO1)3 8H2O, which seem to be given by the sulphates of all the rare earth metals except lanthanum

The densities of the sulphates are given in the following table -

|                                        | La.  | Ce.          | Pr.          | Nd.  | Sm.  | Gd   | Y    | Er.  |
|----------------------------------------|------|--------------|--------------|------|------|------|------|------|
| Anhydrous . 5H2O                       | 3·60 | 3 91<br>3 17 | 3·72<br>3 17 |      |      | 4.14 | 2 61 | 3.68 |
| 8H <sub>2</sub> O<br>9H <sub>2</sub> O | 2.83 | 2 89<br>2 84 | 2 82         | 2 85 | 2 93 | 3.01 | 2 54 | 3 20 |

<sup>1</sup> Chavastelou, Compt. rend., 1900, 130, 781; Giossmann, Zeitsch anorg. Chem., 1905,

<sup>44. 229 (</sup>Ce, La).

Jolin, Bull. Soc. chim., 1874, [ii.], 21, 533 (Ce), Cleve, ibid., 1874, [ii.], 21, 196 (La);
Chem. News, 1886, 53, 81 (Sm), Cleve and Hoglund, Bull. Soc. chim., 1873, [ii.], 18, 193,

In concentrated solutions a slight precipitation is observed in the yttimin group (Crookes, ('hem. News, 1896, 74, 259).

See fig. 39 on p. 383.

With the exception of the cerium salt, all the octahydrates of the sulphates crystallise in monoclinic prisms, isomorphous with one another and with the corresponding selenates:-1

| Salt.                                                             | a. b.      | c.     | β.       |
|-------------------------------------------------------------------|------------|--------|----------|
| Pr <sub>2</sub> (SO <sub>4</sub> ) <sub>8</sub> 8H <sub>2</sub> O | 2 9863 · 1 | 1 9995 | 118° 0′  |
| Nd2(SO4)3.8H2O                                                    | 2.9835 - 1 | 1 9968 | 118° 85′ |
| $Sm_{2}(SO_{4})_{3} 8H_{2}O$                                      | 3 0030 1   | 2 0022 | 118° 16′ |
| $Gd_{2}(SO_{4})_{3}.8H_{2}O$                                      | 3 0086 1   | 2 0068 | 118° 2′  |
| Y,(SO,), 811,0                                                    | 3 0284 1   | 20092  | 118° 25′ |
| Er,(SO <sub>4</sub> ), 811,0                                      | 301201     | 2 0043 | 118° 27′ |

The enneahydrates of cerum and lanthanum crystallise in the hexagonal system (bipyramidal), isomorphous with one another -

$$\begin{array}{lll} \text{La}_2(\text{SO}_4)_3.9\text{H}_2\text{O}, & a & c=1 & 0.7356 \\ \text{Co}_2(\text{SO}_4)_3.9\text{H}_2\text{O}, & a & c=1 & 0.7310 \end{array}$$

The pentahydrates of cerium and prascodymium are also isomorphous with one another (monoclinic prisms) -

The order of solubility of the hydrated sulphates in water at 25° is as follows, beginning with the least soluble -

The actual solubilities at 25°, in grams of anhydrous sulphate per 100 grams of solution, are as follows -2

| Solid Phase                                                                             | Solubility     | Solid Phase.                                                                                      | Solubility.      | Solid Phase                                            | Solubility. |
|-----------------------------------------------------------------------------------------|----------------|---------------------------------------------------------------------------------------------------|------------------|--------------------------------------------------------|-------------|
| $\text{La}_2(\text{SO}_4)_4 9 \text{H}_2 O$                                             |                |                                                                                                   |                  | Pr <sub>2</sub> (SO <sub>4</sub> ), 811 <sub>2</sub> O |             |
| - Gdj(SO <sub>1</sub> ), 8H <sub>2</sub> O<br>- Sm(SO <sub>1</sub> ), 8H <sub>2</sub> O | 2 981<br>3 426 | Nd <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> ,8H <sub>2</sub> 0<br>  Co.(SO.)8H <sub>2</sub> 0 | 0 530  <br>0 760 | $E_{1_2}(SO_4)_3 8\Pi_2O$                              | 1194        |

The solubilities of a number of the preceding salts in aqueous sulphuric acid have been determined at  $25^\circ$  . The results are shown to scale in fig. 20.

Double Sulphates.—The sulphates form numerous double sulphates with the sulphates of ammonium and the alkali metals. Disregarding the water of crystallisation, they may be classified into the following types :-

the water of crystalisation, they may be classified into the following types — 
$$3M_2(SO_4)_3.2R_2SO_4,\ M_2(SO_4)_3.R_2SO_4;\ 2M_2(SO_4)_3.3R_2SO_4,\ M_2(SO_4)_3.3R_2SO_4;\ 2M_2(SO_4)_3.1R_2SO_4,\ 2M_2(SO_4)_3.9R_2SO_4;\ M_2(SO_4)_3.5R_2SO_4.$$

Only one or two systematic investigations of the double sulphates have been made

The double potassium sulphates of the cerium and terbium groups are very sparingly soluble in cold water, and in a cold, saturated solution of

Data from Groth, Chemische Krieballographie (Leiprig, 1906-1910), vol 11
 See Witth, Zeitsch anora, Chem., 1912, 76, 174. James and Holden, J. Amer. Chem.
 Soc., 1913, 35, 559. no data are available for rare earth sulphates other than those quoted.
 Wirth, loc. cit.

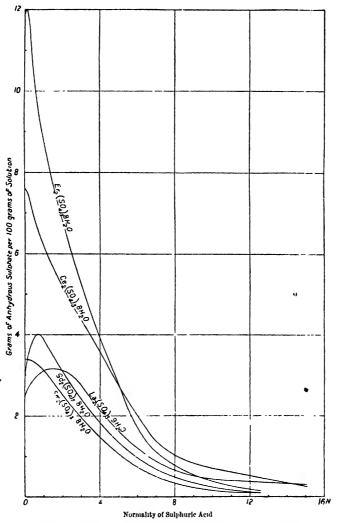


Fig. 20. —Solubilities of the sulphates of some of the rare earth elements in aqueous sulphuric acid at  $25^{\circ}$  C.

potassium sulphate they are even less soluble. In fact, the lanthanum, cerum, prascodymum, and neodymum salts are practically insoluble in the sulphate solution 1 On the other hand, the double potassium sulphates of the yttimm group do not differ greatly from the simple rare earth sulphates in their solubility in water, and are readily soluble in cold, saturated potassium sulphate solution.

The double sodium and ammonium sulphates resemble the potassium salts in their solubilities, but are rather more soluble, particularly the ammonium salts (see pp. 385, 408, 423)

The double ammonium salts of the type M<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub> (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> 8H<sub>2</sub>O form an isomorphous series, crystallising in monochnic prisms —

| Salt. | а      | b | c       | β.      | Dennity |
|-------|--------|---|---------|---------|---------|
| La    | 0 3509 | 1 | 0 9145; | 97° 36′ | 2 5 1 6 |
| C'e   | 0.3598 | 1 | 0.9340  | 97° 15′ | 2.523   |
| "Di"  | 0.3417 | 1 | 0.9234  | 95° 28′ |         |

The double sulphates are of great practical importance. The sparingly soluble double salts may be brought into solution by heating them with concentrated ritric acid and pouring the whole into boiling water, or by dissolving them in concentrated ammonium acctate solution. They may be converted directly into hydroxides by heating with alkali hydroxide,5 and into oxalates by prolonged heating with aqueous oxalic acid By heating to redness for an hour or so with four times their weight of powdered charcoal, they are almost completely reduced to sulphides, which may be dissolved in hydrochloric acid 7

Acid Sulphates, M(HSO<sub>4</sub>), -These salts may be prepared by dissolving the nomal sulphates in boiling sulphuric acid and cooling the solution, when the crystalline acid salts separate out, " or by dissolving the normal salts in ice cold water and adding a large excess of sulphune acid.9 Adhering sulphuric acid may be removed by heating in vacuo to 130°, but above 180° decomposition commences.

The cerous salt forms colourless, glistening needles, the yttimm salt, coloniless pyramids; the lanthanum, prascodymium, neodymium, and samarının salts form colourless, green, rose red, and golden-yellow, sılky needles respectively. All the acid salts are converted into normal salts at temperatures approaching dull redness

Basic Sulphates, M2O3SO3 -- It is only at a white heat that a rare earth sulphate can be completely converted into the oxide. When heated to 900°,

<sup>1</sup> This statement does not rigidly apply when mixtures of rare earth sulphates are dealt

with, see p. 338

2 Morton, Zettsch. Kryst. Min., 1887, 12, 520. Krans, thid., 1901, 34, 307. Wyrouboff,
Bull. Soc. Trang. Vin., 1891, 14, 83.

3 Hranger and Berzelne, Ann. Chim., 1803, [1], 50, 245, R. J. Meyer and Marckwald,

Ber., 1900, 33, 3003.

4 Urbain, Bull. Soc. chim., 1496, [m], 15, 338, 347, of Muthmann and Rohg, Ber.,

<sup>Urbain, Itall. Soc. chim., 1896, [iii], 15, 338, 347, cf. Muthinain and Rong, Ber., 1898, 31, 1718
Von Scheele, Ber., 1899, 32, 409.
Auer von Welshach, Monatsh., 1883, 4, 630. Muthinain and Rong, loc cd.
Browning and Blumenthal, Amer. J. Ser., 1911, [iv], 32, 161, Dennis and Rhodes, J. Amer. Chem. Soc., 1915, 37, 809.
Wyrouboff, Bull. Soc. chim., 1889, [iii], 2, 745 (Ce); Matignon, Compt. rend., 1902, 134, 657 (Pt., Nd.); 1905, 141, 1230 (Sm.)
Brauner and Picek, Zettsch. anory. Chem., 1904, 38, 322 (Ls., Ce, Pr., Nd., Sm., Y).</sup> 

basic sulphates of the type M2O3 SO3 are obtained (except from cerous sulphate) as amorphous powders, stable at 1000° 1

Selenites, M<sub>a</sub>(SeO<sub>a</sub>),—The precipitates obtained by adding solution selenite to an aqueous solution of rare earth salts are usually basic selenites. When a solution of sclemous acid is used as precipitant, the normal sclemes are generally precipitated. By heating the basic scienitis with dilute aqueous selenious acid, the normal scienites may be obtained, and if more concentrated selemous acid be employed, acid selemites are formed 2

Selenates,  $M_2(SeO_4)_3$ —The sclenates may be obtained as hydrated salts by dissolving the oxides, hydroxides, or carbonates in aqueous sclenic acid and crystallising from the solution. By carefully heating the hydrated salts, anhydrous selenates may be obtained. The selenates are not so well known as the sulphates A number of octahydrates are known, and those that have been measured are isomorphous with the corresponding sulphates -3

The decallydrates of the same three elements crystallise in orthorhombic bipyramids and are isomorphous with one another, the data for two are as follows --

The densities of the selenates are given in the following table --

|                                                             |          |     |      |      |      | -    |      | - |
|-------------------------------------------------------------|----------|-----|------|------|------|------|------|---|
|                                                             |          |     | La   | P1.  | Sm   | Gd   | • Y  |   |
|                                                             |          |     |      |      |      |      | ļ    | ٠ |
| Anhydrous                                                   | ,        |     |      | 4 30 | 4.08 | 4 18 |      |   |
| 6H <sub>2</sub> O<br>8H <sub>2</sub> O<br>9H <sub>2</sub> O | :        | • ; | 3.48 | 3:08 | 3 33 | 3 32 | 2 91 |   |
| 10H <sub>2</sub> O<br>12H <sub>2</sub> O                    | :        |     | 2*66 |      | 3 01 | 3 06 | 2.78 |   |
| 121190                                                      | <u>.</u> |     | -    |      |      | ł    |      |   |

In general, the double selenates are rather more soluble in water than the corresponding double sulphates.

Chromates, M2(CrO1)3 — Few of these salts appear to have been analysed. The chromates are sparingly soluble in water, and so may be obtained as crystalline precipitates by double decomposition, using potassium chromate as precipitant. When a large excess of precipitant is used, double chromates may be obtained, particularly in the yttium group. The chromates appear as a rule to have the composition M2(CrO4), 81120 and to be isomorphous with the corresponding sulphates.

<sup>&</sup>lt;sup>1</sup> Matignon, Compt. rend., 1902, 134, 657 (Nd, Pr); 1905, 141, 1230 (Sm), 1906, 142, 394; Wild, Zeitsch. anorg. Chem., 1905, 47, 464 (La. Sm, Y. E., "Yb"), L. Wohler and Grunzweig, Ber., 1913, 46, 1726 (La. Pr., Nd, Sm, Gd. Y. E., Yb, Lu).

<sup>8</sup> Nilson, Nova Aida Noc Upsala, 1875, [m], 9, No 7; Ber., 1875, 8, 655; Bull. Soc. chim., 1875, [i.], 23, 494

<sup>3</sup> Topsoc, Bihang K. Sienska Vet Akad. Handl., 1874, 2, No. 5; Bull. Soc. chim., 1874, [n], 22, 353; Ber., 1875, 8, 129 (E1, Y). Benedicks, Zeitsch. anorg. Chem., 1900, 22, 598 (Gd).

...

The rare earth salts are not precipitated by the addition of chromic acid or an alkalı dichromate, in this respect they differ from those of zirconium, thorium, and quadrivalent cerium

The chromates are of value for the separation of the rare earths from

one another (see pp 343, 352).

Molybdates, M. (MoO<sub>4</sub>)<sub>3</sub>.—These salts closely resemble the corresponding tungstates in crystalline form and properties, and may be prepared by means similar to those described in the next section for the preparation of tungstates. Cerous and "didymium" molybdates have been shown by Cossa to be isomorphous with lead molybdate 1

A number of complex molyblates have been prepared containing the rare earth elements of the cornum group. They have the general formula (NH<sub>4</sub>)<sub>0</sub>M<sub>2</sub>Mo<sub>14</sub>O<sub>15</sub> 24H<sub>2</sub>O<sub>5</sub> and form a series of isomorphous salts crystallising in the triclinic system -2

|               | a i        | c      | a        | В       | γ        |
|---------------|------------|--------|----------|---------|----------|
| La            | 0 3502 1   | 0.3416 | 102° 29′ | 51° 18' | 103° 10′ |
| Co            | 0.3523 - 1 | 0.3409 | 102° 22′ | 54° 30′ | 103° 4'  |
| $\mathbf{Pr}$ | 0 3511     | 0 3461 | 102* 11′ | 54° 15′ | 103° 45′ |
| Nd            | 0 3492 1   | 0.3385 | 102° 15′ | 54° 8′  | 103° 29′ |
| Sm            | 0 3611 1   | 0.3330 | 102° 49′ | 54 15'  | 102° 37′ |

Tungstates,  $M_2(WO_4)_{\gamma}$ .—The normal tungstates of the rare earth elements are practically insoluble in water, and may be obtained by double decomposition between sodium tungstate and a rare earth intrate or sulphate The precipitates are usually amorphous, but may become crystalline when heated. The amorphous tungstates may be crystallised by fusing them in an mert atmosphere and cooling, or by mixing them with an excess of sodium chloride and fusing the mixture. The tungstates crystallise in tetragonal bipyramids It was discovered by Cossa that cerous, "didyneum," and calcium tungstates are isomorphous and form mixed crystals, and Zambonini has shown that cerous and lead tungstates are completely miscible both in the liquid and the solid states

When a rare earth is fused with tungstic acid, sodium tungstate, and sodium chloride, and maintained at a red heat for some time, double tungstates are formed and may be isolated by washing the mass with water, in which they are insoluble. The following salts have been thus obtained in crystals which closely resemble those of the simple rare carth tungstates - 5

| La <sub>4</sub> Na <sub>6</sub> (WO <sub>4</sub> ) <sub>9</sub> | $\operatorname{Sm}_{4}\operatorname{Na}_{6}(\operatorname{WO}_{1})_{s}$ |
|-----------------------------------------------------------------|-------------------------------------------------------------------------|
| La, Na, (WO),                                                   | Er <sub>4</sub> Na <sub>6</sub> (WO <sub>1</sub> ) <sub>9</sub>         |
| Ce <sub>2</sub> Na <sub>8</sub> (WO <sub>1</sub> ) <sub>7</sub> | Y2Na8(WO1)7                                                             |

<sup>&</sup>lt;sup>1</sup> Frenchs and Smith, Annalen, 1878, 191, 331 (La<sub>2</sub>(HMoO<sub>4</sub>)<sub>2</sub>), Cossa, Compt. rend., 1884, 98, 990 ("Di"); 1886, 102, 1315 (Ce), Gazz tta, 1886, 16, 284 ('e, "Ih"), Cleve, Chem News, 1886, 53, 93 (Sm), Hitchcock, J. Amer. Chem. Soc., 1895, 17, 520 (La, Ce,

The metatungstates of the rare earth metals are extremely soluble in water, and may be prepared from baruum metatungstate and the sulphates The lanthanum, cerum, and "dadymium" salts form two series of hydrates,  $M_2(W_1O_{1a}), 271I_2O$  and  $M_2(W_4O_{13}), 301I_2O$ , the latter crystallising below 16 in the triclinic, the former above 16 in the orthorhombic system —

```
abc
                             30H<sub>2</sub>O.
  a b : c.
   97° 6′ 93° 50′ 89° 2′
96° 13′ 92° 34′ 91° 30′
"Di" 0 9736 1 . 1 2952 La 0 9859 1 0 5786
Ce 0 9838 1 0 5825
```

A samarum salt, Sa<sub>2</sub>(W<sub>1</sub>O<sub>13</sub>)<sub>3</sub>.35H<sub>2</sub>O, has also been described.<sup>1</sup>

A number of very complex tungstates, in which rare earth and tungstic anhydrale are in the ratio M2O3. 16WO, have been prepared by Rogers and E. F. Smith.<sup>2</sup> The formulæ of these compounds are given below

```
\begin{array}{c} 2(\mathrm{NII_{1}})_{2}\mathrm{O}\ \mathrm{La_{2}O_{3}}\ 16\mathrm{WO_{3}}\ 16\mathrm{H_{2}O} \\ 5\mathrm{BaO}.\ \mathrm{La_{3}O_{3}}\ 16\mathrm{WO_{3}}\ 16\mathrm{H_{2}O} \\ 5\mathrm{Ag_{2}O}\ \mathrm{La_{3}O_{3}}\ 16\mathrm{WO_{3}}\ 4\mathrm{H_{2}O} \\ 3(\mathrm{NII_{1}})_{2}\mathrm{O}\ \mathrm{Nd_{2}O_{3}}\ 16\mathrm{WO_{3}}\ 4\mathrm{H_{2}O} \\ 3(\mathrm{NII_{1}})_{3}\mathrm{O}\ \mathrm{Nd_{2}O_{3}}\ 16\mathrm{WO_{3}}\ 4\mathrm{H_{2}O} \end{array}
\begin{array}{c} 2(\mathrm{NII_4})_2\mathrm{O}.\mathrm{Ce_2O_3}\ 16\mathrm{WO_3}.2\mathrm{H_2O} \\ 2(\mathrm{NII_4})_2\mathrm{O}.\mathrm{Pr_2O_3}.16\mathrm{WO_3}.16\mathrm{H_2O} \\ 4\mathrm{BaO}\ \mathrm{Pr_2O_3}.16\mathrm{WO_3}.7\mathrm{H_2O} \\ 2\mathrm{Property}. \end{array}
   6BaO.Pr<sub>2</sub>O<sub>3</sub>.16WO<sub>3</sub> 9H<sub>2</sub>O
   4Ag<sub>2</sub>O Pr<sub>2</sub>Ö<sub>8</sub>.16WÖ<sub>3</sub> 8IĨ<sub>2</sub>O
  6BaO, Nd<sub>2</sub>O<sub>5</sub>, 16WO<sub>2</sub>, 17H<sub>2</sub>O
```

Silicotung states,  $M_i(W_{12}SiO_{10})_{ij}$ , i  $\epsilon$ .  $2M_2O_3$  [12 $WO_3$   $SiO_2$ ]<sub>3</sub> —The normal and acid silicotungstates of the rare earth elements are soluble salts which orystallise well. They have been examined by Wyrouboff's The following types of normal salts are known :-

(i)  $M_1(W_{12}S(O_{10}), 78H_2O,$  crystallising in the trigonal system (Nd, Sm, Gd, Th, Y, "Yb" salts known). The ratio ca (vertical to lateral crystallographic axis) is as follows -

```
Salt.
      Nd
             8m
                      Gd.
                              Tb.
                                      Y.
    1.6636 1.7244 1.7144 1.7043 1.7120 1.69.)7
c/a
```

(ii ) M<sub>4</sub>(W<sub>1</sub>,SiO<sub>40</sub>)<sub>3</sub> 8HH<sub>2</sub>O, crystallising in the trigonal system (Lo, Ce, Pr, Nd, Sm, Gd salts known) --

```
Salt.
                          "Di"
      26392
                2 6820
                          26653
                                   26786
   26653
c/a
```

The cerium and "didymium" salts are also known in monochine forms .-

```
a = b = 17090 = 1 = 26558.
  \beta = 90' 20'
"Di"
  B== 90° 10'
           a b c = 17000 1 26325,
```

(m.)  $M_4(W_{12}S_1O_{10})_4$  90H<sub>2</sub>O —Only the gadolinium salt is known (trigonal;  $a \cdot c = 1 + 2.6238$ ).

The acid salts also fall into three classes -

(1) MH( $W_{12}S1O_{40}$ ) 18H  $_2O$ , crystallising in the trichine system (La, Ce, Pr, Nd, Sm, Gd salts known) —

| Salt                                                                | $a \cdot b \cdot c$ . | α.      | β.      | γ.      |
|---------------------------------------------------------------------|-----------------------|---------|---------|---------|
| $LaH(W_{12}S_1O_{40}) 18H_2O$                                       | 0.4081 1 . 0.4420     | 89° 34′ | 93° 8′  | 81° 29' |
| $C_0H(W_{13}S_1O_{40})$ 18H,0                                       | 0.3922 1 0 4195       | 90° 43′ | 92° 20' | 84° 38′ |
| " Di " $\dot{H}(\dot{W}_{12}S_1\ddot{O}_{40}).18\ddot{H}_2\ddot{O}$ | 0.1003 1 0.4262       | 89° 58  | 93, 48, | 83° 20′ |
| GdH(W <sub>12</sub> S1O <sub>40</sub> ).18H <sub>2</sub> O          | 0.4127 . 1 0.4393     | 89° 20′ | 92° 35′ | 84° 9′  |

Scheibler, J. prakt Chem., 1861, 83, 273 (Ce), Wyrouboff, Bull. Soc franç. Min.,
 1892, 15, 63 (La, Ce, "In"), Cleva Chem. News, 1886, 53, 93 (Sm)
 Rogers and Smith, J. Amer. Chem. Soc., 1904, 26, 1474.
 Wyrouboff, Bull. Soc franç Min., 1896, 19, 219, 1905, 28, 201.

(ii.) MH (W $_{12}{\rm SiO}_{40})$  24  $_2^{1}{\rm H}_2{\rm O}$  . Monoclinic crystals (Tb, Y, Er, "Yb" salts known) --

| Salt.                                        | a b c           | β.     |
|----------------------------------------------|-----------------|--------|
| $TbH(W_{12}S_1O_{40}) 24\frac{1}{2}H_2O$     | 1 0270 1 1.4132 | 92° 39 |
| YH(Ŵ <sub>a</sub> ŠiO <sub>t</sub> o)~24JH.Ō | 1.0446 1 1 4835 | 92° 4  |
| "Yb `H(W, SiO <sub>10</sub> ) 24JH,O         | 1 0356 1 1 1747 | 92° 5  |

(iii ) MH(W<sub>12</sub>SiO<sub>40</sub>) 25½H<sub>2</sub>O.—Only the *orbium* salt is known (trigonal; a e=1 17244)

It is an interesting fact that the normal silicoting states of lanthanum, cerum, and "didymium" with 81H<sub>2</sub>O are isomorphous with thorum silicoting state and the silicoting states of calcium and strontium (see p. 234).

#### THE RARE EARTH ELEMENTS AND THE NITROGEN GROUP.

Nitrides, MN—The fact that the rare earth metals combine directly with introgen to produce intrides was discovered by Matignon. The compounds are of the type M<sup>III</sup>N, as Muthinain, Kraft, and Beck have shown, and are readily prepared by heating the metals to 900° m introgen. Moissan has demonstrated the formation of intrides when the carbides are heated to 1000′ m ammonia, and Datert and Miklanz¹ have shown that ecrons intride may be obtained in a pure condition by heating the hydride to 800°-900° m pure nitrogen.

The intrides are brittle, amorphous solids, the cerum compound being biass-yellow and the compounds of lanthanum, pracedymium, and needymium black in colour. They are decomposed by moist air or by water, hydroxides and ammonia being produced. The atomic heat of introgen in the cerum and landbanum compounds is 4.9.1

Nitrites, M(NO<sub>2</sub>). - Practically nothing is known of the rare earth intrites. A number of phitinonitrites and platino-iodonitrites have been prepared by Nilson.<sup>1</sup>

Nitrates, M(NO<sub>3</sub>)<sub>p</sub>—These salts are readily obtained in solution by dissolving the sesqui-oxides, hydroxides, or carbonates in intric acid, or by oxidising the oxidates with hot, concentrated intric acid. They are readily soluble in water, from which the majority crystallise as hexahydrates, M(NO<sub>2</sub>)<sub>3</sub>6H<sub>2</sub>O. They are readily soluble in alcohol and also in acctone.<sup>5</sup> When heated they decompose, giving rise to basic salts and finally to the oxides.

The solubilities of a number of the hevally drates in water at 25 are as follows (in parts of anly drons intrate per 100 of water) =  $^6$ 

These chemists deny that cerous nitide may be prepared by heating common miningen.
 Matignon, Computered, 1900, 131, 837 (La. Ce. Pr. Nd, Sm); Moissan, vbid, 1900, 131, 595, 865 (Pr. Nd), Muthmann and Kialt, Annation, 1902, 325, 261 (Cc. La); Mithmann and Beck, vbid., 1904, 331, 58 (Pr. Nd), Dafert and Mikhaur, Monatch, 1912, 22, 911 (Cc.)

Kellenberger and Kraft, Annalen, 1902, 325, 279 Temperature interval not stated,
 Nilson, Nova Acta Soc Upsala, 1879. [m], 10, No. 16, Eer., 1876, 9, 1722; 1878,

<sup>11, 879

&</sup>lt;sup>5</sup> Barnebey, *J. Amer. Chem. Soc.*, 1912, 34, 1174.

<sup>6</sup> James and Pratt, *J. Amer. Chem. Soc.*, 1910, 32, 873 (V), James and Robinson, *ibid.*, 1913, 35, 754 (Nd); James and Whittemore, *ibid.*, 1912, 34, 1168 (La).

The solubilities are diminished by the addition of nitric acid, but from concontrated nitric acid (dens 1.4) the pentahydrates, M(NO<sub>3</sub>)<sub>3</sub> 5H<sub>2</sub>O, usually separate out The fractional crystallisation of the mitrates is often used in separating the rare earths

The hexahydrates crystallise in the triclinic system, as the following results 1 indicate -

|                                       | авс                       | a       | β        | γ        |
|---------------------------------------|---------------------------|---------|----------|----------|
| La(NO <sub>2</sub> )6H <sub>2</sub> O | 0 8315 1 0.6215           | 79° 12′ | 101° 58' | 92° 18′  |
| $Ce(NO_3)_3.6H_2O$                    | $0.8346 - 1 \cdot 0.6242$ | 78° 51  | 102 9'   | 92° 3′   |
| "Di(NO <sub>3</sub> )",611,0 "        | 2.4035 1 1 8597           | 69, 6,  | 97° 30′  | 92° 48′  |
| Gd(NOL) 61LO                          | 0 5571   0 7615           | 90° 6′  | 109° 55′ | 109° 48′ |

The nitrates of needymnum, lanthanum, and yttinum have been shown to be isodimorphous with bismuth nitrate (p. 231)

Since double intrates are rather uncommon, it is interesting to notice that the nitrates of the rare earth metals form numerous double salts, of great practical value The nitrates of the metals of the cerum group form double ammonium initiates of the type 2(NH<sub>4</sub>)NO<sub>3</sub> M(NO<sub>3</sub>)<sub>3</sub> HI<sub>2</sub>O, crystallising in monochmic prisms.2

```
\beta = 112^{\circ} 36'

\beta = 112^{\circ} 45'

\beta = 113^{\circ} 1'
                                a.b c- 1.2475 1 2.1863,
                                a \ b \ c = 1 \ 2321 \ 1 \ 2 \ 1695
```

Of the double nitrates formed with alkali nitrates, only those containing rubidium are of the type 2MNO, X(NO<sub>3</sub>), III<sub>3</sub>O, and isomorphous with the ammonium salts The following data are due to Wyrouboff

```
101° 16′
  1:2298 1 0 9910
2RbNO, Ce(NO<sub>3</sub>), 4H<sub>2</sub>O
   101°
2RbNO_{3}^{2}La(NO_{3}^{3})_{3}^{4}4H_{2}^{2}O
2RbNO_{3}^{2}Nd(NO_{3}^{3})_{3}^{4}H_{2}O
  1 2319 1 0 9888
   100° 29′
  1 2276 1 . 1 0153
```

The corresponding prascodymium-rubidium salt has been prepared by Jantsch and Wigdorow,4 who have likewise prepared the thallium double salts

$$2TINO_3.Ce(NO_3)_3.4II_2O \quad and \quad 2TINO_3.La(NO_3)_3.4H_2O,$$

which they state are isomorphous with the preceding salts

The following melting-points and densities of the preceding double intrates have been determined, mainly by Jantsch and Wigdorow -

|                             | Ce.                 | l.a.       | Pr.   | Nd      | Ce.            | La             | Pr.           | Nd   |
|-----------------------------|---------------------|------------|-------|---------|----------------|----------------|---------------|------|
| NH <sub>4</sub><br>Rb<br>Tl | 74°<br>70°<br>61.5° | 86°<br>72° | 63 5° | <br>47° | 2·497<br>3 326 | 2·497<br>3·318 | 2 151<br>2·50 | 2.56 |

apply.

Jantsch and Wigdorow, Zeitsch anorg. Chem., 1911, 69, 221.

<sup>&</sup>lt;sup>1</sup> Taken from Groth, Chemische Krystallographie (Leipzig, vol. 11, 1908)
<sup>2</sup> Data from Groth, opus cit., cf. Wyrouboff, Bull. Soc franc. Min., 1906, 29, 324.
<sup>3</sup> Wyrouboff, Bull. Soc. franc. Min., 1907, 30, 299, the choice of axes and parametral plane is not the same as that for which the preceding data concerning ammonium salts

"Ý

The other double nitrates containing the alkali metals will be described under the headings of the rare earth metals. All the double intrates are best crystallised from fairly concentrated mitric acid at comparatively low temperatures, e.g. 30° C. At temperatures above 70°, anhydrous double salts separate out <sup>1</sup> They are of the type 2M<sup>11</sup>(NO<sub>4</sub>), 3M<sup>1</sup>NO<sub>3</sub>, crystallise in the cubic system, and rotate the plane of polarised light <sup>2</sup> When kept at the ordinary temperature, they become converted into other polymorphic forms having a low degree of symmetry a

With the intrates of magnesium, nickel, cobalt, sine, and manganese, the mitrates of the cerum and terbuna metals form double salts of the type 2M<sup>III</sup>(NO<sub>3</sub>), 3M<sup>II</sup>(NO<sub>3</sub>), 24H<sub>3</sub>O. These double intrates form a comprehensive group of isomorphous salts, crystallising in the trigonal (scalenohedral) system, isomorphous with the corresponding bismuth double salts. The following values for the ratio (c) of the vertical to the lateral axes have been recorded -- 1

|                     | Mg                | Mu     | Ni     | Co     | Zn     | Fe (ous). |
|---------------------|-------------------|--------|--------|--------|--------|-----------|
| ('e<br>Nd<br>'' Dı" | 1 5 :78<br>1 5720 | 1 5775 | 1 5667 | 1.5742 | 1 5077 | 1 5590    |
| Gď                  | 1:5786            |        |        | 1.     | 1      | 1 5590    |

The double salts melt in their water of crystallisation at the following temperatures: -- 5

|    | Mg.                                              | Mu                                  | Nı.                                               | ( 0                                           | Zn                                   |
|----|--------------------------------------------------|-------------------------------------|---------------------------------------------------|-----------------------------------------------|--------------------------------------|
| La | 113°5<br>111 5<br>111 2<br>109 0<br>96 2<br>77°5 | 47°2<br>>37<br>81°0<br>77°0<br>70°2 | 110°5<br>108 5<br>108 0<br>105 6<br>102 2<br>72 5 | 111 8<br>98 5<br>97 0<br>95 5<br>83 2<br>63 2 | 95°0<br>92 8<br>91 5<br>88 5<br>76 5 |

The order of fusibility is therefore the same in each series.

The solubilities of the double intrates in intric acid (of density 1 325 at 16°) have been determined by Jantsch. The following data represent the solubilities in gram molecules of salt per litre at 16°

|     | ·     | Иg     | Mn.    | Nı     | Co.    | , Zu   |
|-----|-------|--------|--------|--------|--------|--------|
|     | <br>  |        |        |        |        |        |
| Ce  | 1     | 0.0382 | 0 1103 | 0.0100 | 0 0632 | 0 0675 |
| l,a | . '   | 0 v418 | 0.1192 | 0 0492 | 0.0569 | 0 0751 |
| Pr  |       | 0 0503 | 0 1442 | 0 0568 | 0 0794 | 0.0858 |
| Nd  |       | 0 0635 | 0 1816 | 0 0710 | 0 0923 | 0 1066 |
| Sm  | - 1   | 0.1583 | 0 3017 | 0 1760 | 0.2072 | 0 2179 |
| Gd  |       | 0 2252 |        | 0 2405 | 0 2706 | 0 2801 |
|     | - 1   |        | 0 3712 | 0 2612 | 0 3090 | 0 3215 |
| Bı  | <br>ı | 0 2503 | 0 3712 | 0 2612 | 0 3090 |        |

<sup>&</sup>lt;sup>1</sup> Except in the case of the sodium salts (or casum? the original paper is ambiguous).

<sup>2</sup> Class 28 of Vol. I. p. 53.

<sup>3</sup> Wyroubolf, Bull. Soc. frant. Man., 1907, 30, 249, 1909, 32, 365.

<sup>4</sup> Gerpel, Zeitsch. Kryst. Min., 1902, 35, 625, Fock, ibid., 1894, 22, 37; Kraus, ibid., 1901, 34, 430; Groth, Chemische Krystallographie (Leipzig, 1906-1910), vol. ii.

<sup>5</sup> Jantsch, Zeitsch. anorg. Chem., 1912, 76, 303

The densities at  $0^{\circ}$  C and the gram-molecular volumes of the double nitrates are as follows, the data being also due to Jantsch.—

|                                  | Mg.                                                |                                                    | Mn                                        |                                           | N                                                  | Nı.                                                 |                                                    | Co                                                 |                                                    | n,                                                 |
|----------------------------------|----------------------------------------------------|----------------------------------------------------|-------------------------------------------|-------------------------------------------|----------------------------------------------------|-----------------------------------------------------|----------------------------------------------------|----------------------------------------------------|----------------------------------------------------|----------------------------------------------------|
| La<br>Ce<br>Pr<br>Nd<br>Sm<br>Gd | 1 988<br>2 002<br>2 019<br>2 020<br>2 088<br>2 163 | 763:3<br>764:2<br>758:0<br>761:2<br>742:4<br>728:0 | 2 080<br>2 102<br>2 109<br>2 114<br>2 188 | 778 6<br>771 6<br>769 3<br>771 0<br>750 3 | 2 116<br>2 173<br>2 195<br>2 202<br>2 272<br>2 356 | 7.59·7<br>751·5<br>744·3<br>745·3<br>727·7<br>707·0 | 2 131<br>2:157<br>2 176<br>2 195<br>2 237<br>2 315 | 765 5<br>757 5<br>751 1<br>748 0<br>739 2<br>720 5 | 2·161<br>2·1·8<br>2·215<br>2·208<br>2·283<br>2·351 | 763 8<br>755 5<br>751 0<br>750 0<br>732 8<br>717 5 |

The more soluble of the preceding double intrates are best crystallised from nitric acid.

The intrates of the metals of the yttrium group do not appear to combine with other intrates  $^{\rm 1}$ 

Orthophosphates, MPO<sub>e</sub>.—Few phosphates of the rare earth elements have been studied in any detail, such information as is available will be found under the headings of the various metals. Certain phosphates occur naturally, e.g. cerous phosphate or monazite (p. 218) and yttrium phosphate or xenotime (p. 220).

When orthophosphoric acid or an alkali orthophosphate is added to a solution of a rare carth salt a gelatinous precipitate of an orthophosphate is produced, which becomes crystalline on standing. The precipitate is soluble in excess of orthophosphoric acid, but is reprecipitated when the solution is boiled; it is also soluble in dilute mineral acids.

## THE RARE EARTH ELEMENTS AND THE CARBON GROUP

Carbides, MC<sub>2</sub> - The carbides of the rare earth metals are readily prepared by heating intimate mixtures of the rare earths and sugar carbon in the electric furnace —

$$M_2O_a + 7C = 2MC_2 + 3CO$$

They were first prepared by Pettersson, and have been examined in detail by Moissan.<sup>2</sup> It will be noticed that the carbides of the rare earth elements are of the same type as calcium carbide, CaC<sub>22</sub> and are not analogous to aluminium carbide. ALC.

The carbides are brittle, crystalline solids, which in thin layers are transparent and yellow in colour. Their densities are as follows --

The carbides burn in fluorine when warmed, giving rise to fluorides and carbon. They likewise burn in chlorine, bronnine, and rodine vapour at

<sup>&</sup>lt;sup>1</sup> For double compounds with organic substances, see Kolb, Zeds h anorg Chem., 1908, 60, 123, 1913, 83, 143; Barbieri and Calzolaii, Alli R. Accad Lincei, 1911, [v], 20,

 <sup>50, 125, 1915, 63, 145,</sup> Instituted and Calzonal, Act. M. Act. B. Matter Editor, 1917, [1], [2], i. 164
 Pettersson, Ber., 1895, 28, 2419 (Ce, Ls); Moissan, Compt. rend., 1896, 122, 357 (Ce), 123, 148 (Ls); 1900, 131, 595 (Pr, Nd), 924 (Sm); Moissan and Etard, ibid., 1896, 122, 578 (Y), Moissan, Ann. Cham. Phys., 1896, [vii.], 9, 302 (Ls, Ce); 1901, [vii.], 22, 110 (Sm), Muthmann, Hofer, and Weiss, Annalen, 1902, 320, 231 (Ce).

temperatures considerably below a red heat, and yield chlorides or iodides, hydrogen and carbon when heated in a stream of hydrogen chloride or iodide. At a red heat they are decomposed by hydrogen sulphide, and buin readily when heated in a stream of oxygen. They are decomposed by sulphur vapour, but only slowly, selemum at a red heat has more action than sulphur on the carbides. At 1200° they are decomposed by introgen (or better, ammoma), intrides being formed. The carbides are unattacked by cold, concentrated intric and sulphuric acids, but the latter acid oxidises them slowly when heated. They are decomposed by fusion with potassium hydroxide, carbonate, intrate, chlorate, or permanganate

The most interesting property of the rare earth carbides is their behaviour towards cold water, which rapidly decomposes them with the evolution of gas, the formation of a little liquid (unsaturated hydrocarbons), and the precipitation of hydroxides, M(OH), <sup>1</sup> The chief constituent of the gas evolved is acetylene (68-72 per cent ), the constituents next in quantity are hydrogen (5-14 per cent) and ethane (8-13 per cent). Both the acetylene and the ethane are accompanied by small amounts of their homologues. Ethylene is also present (5-9 per cent ), but methane is absent. Apparently the initial reaction is -

$$MC_2 + 3H_2O = M(OH)_3 + C_2H_2 + H_2$$

followed by the hydrogenation of part of the activene to ethylene and ethane 2

Carbonates, Ma(CO<sub>3</sub>), -- That the rare earths are strongly basic oxides is indicated by the fact that normal carbonates of the rare earth elements are easily prepared. They may be obtained in the crystalline form by passing a current of carbon dioxide through aqueous suspensions of the hydroxides, and as amorphous or crystalline precipitates by adding a very dilute solution of an alkali carbonate or bicarbonate to dilute solutions of rare earth salts

The normal carbonates are insoluble in water. When the precipitated carbonates are allowed to stand in contact with concentrated alkali carbonate solutions, they are transformed into crystalline double carbonates. The double salts may be directly precipitated by adding a concentrated solution of a rare earth salt, drop by drop, to a cold, concentrated solution of an alkali carbonate, (1) The double salts of the cerum group of metals are slightly soluble in concentrated sodium or ammonium carbonate, but readily soluble in concentrated potassium carbonate solution, the order of increasing solubility being lanthanum, praseodymium, cerium, and neodymium ! The potassium salts may therefore be prepared by adding the requisite rare carth chlorides to excess of concentrated potassium carbonate solution, warming to dissolve the precipitates, cooling, and diluting slightly, the potassium neodymnum and potassium-prascodymium salts may be obtained in glittering needles by the slow evaporation of their solutions in potassium carbonate (i) The double salts of the yttrum group of elements are much more readily soluble in sodium or ammonium carbonate than those of the cernin group, particularly when warmed 4 All the double carbonates are decomposed

<sup>1</sup> Even in the case of cerum.

Even in the case of centum.
 Dannens, Compt. rend., 1913, 157, 214, cf. the previous references
 Hiller, Inaugural Dissertation (Weilin, 1901), R. J. Meyer, Zeits h. amorg. Chem.,
 1904, 41, 97. Arnold, Ber., 1905, 35, 1173
 Drosslach, Ber., 1900, 33, 3500. Dennis and Dales, J. Amer. Chem. Soc., 1902, 24,
 Arnold, loc. ct.

by water or dilute alkalı carbonates, particularly the sodium salts, and the salts of the cerium group are more readily decomposed than those of the yttrium group.

The sodium, potassium, and ammonium salts are of the type RicOg. M2(CO3)3 xH2O, and sodium salts of the type 3Na2CO3 2M2(CO3)3 xH2O are also known.

Cyanides.—The simple cyanides of the rare earth metals are not known, hydroxides being precipitated when attempts are made to prepare the salts from potassium cyanide and rare earth salt solutions The platinocyanides, however, are beautiful crystalline salts. Those of the metals of the cerium group are monoclinic, yellow in colour with a blue reflex, and belong to the type Pt3M2(CN)12 18H2O, those of the metals of the terbium and yttrium groups are rhombic, red in colour with a green reflex, and belong to the type Pt<sub>8</sub>M<sub>2</sub>(CN)<sub>12</sub> 21II<sub>2</sub>O:—1

|                                                                         | 11     | b | ι.     | β.         | Density |
|-------------------------------------------------------------------------|--------|---|--------|------------|---------|
| $\text{La}_{2}\text{Pt}_{3}(\text{CN})_{12} 1811_{2}\text{O}$           |        |   |        |            | 2626    |
| Ce <sub>0</sub> Pt <sub>1</sub> (CN) <sub>10</sub> .1811 <sub>1</sub> O | 0.5806 | l | 0.5527 | 107" 33'   | 2.657   |
| " $D_{1_{2}}$ " $P_{t_{3}}$ ( $CN)_{1_{2}}$ 1 $81_{2}$ 0                | 0.5806 | l | 0 5517 | 107° 29 5′ |         |

Thiocyanates,  $M(CNS)_3$ .—These salts are readily soluble in water and alcohol. They form double salts with mercuric eyanide, of the type  $M(CNS)_3$   $3Hg(CN)_2$ . $12H_2O$  The lanthanum and cerum salts are monoclinie -

La, a b c=22787 1 25787, 
$$\beta = 92^{\circ} 37'$$
  
Ce, a b c=22921 1 25655,  $\beta = 92^{\circ} 39'$ 

but the yttrum and erbium salts are triclinic -2

Formates, (II COO), M.—The formates of the rare earth elements form well-defined crystals They belong to the regular system, crystallising in pentagonal dodecahedra.3 The formates of the cornim metals are sparingly soluble and those of the yttrium group fairly readily soluble in water, the formates of the terbium metals being intermediate in solubility. These solubility differences have at times been utilised for fractionating the rare earths (see p 317).

Acetates, (CH3 COO),M.—These salts are readily soluble in water. Basic salts do not separate out from dilute boiling aqueous solutions

Topsoe, Bihang K. Svenska Vrt -Akud Handl., 1874, 2, No. 5, Soderstrom, Zeitsch.
 Kryst. Min., 1902, 36, 194 (Pr), Baumhauer, Zeitsch Kryst Min., 1907, 43, 356 (Y);
 Tschrivinski, ibid., 1913, 52, 44 (Y), Biguilawski, Ann Physik, 1914, [iv], 44, 1077 (Y),
 Topsoe, Bihang K. Svenska Vet. Akad. Handl., 1874, 2, No. 5.
 Behrens, Arch. Néerland., 1901, [ii], 6, 67, Rec. trav. chim., 1904, 23, 413.

Oxalates, Mg(C2O4), -The oxalates are readily obtained by the addition of an aqueous solution of ovalic acid to solutions of soluble salts of the rare earth elements, they are perhaps best obtained from the intrates  $^{1-}$  The oxalates are first precipitated in the amorphous state, but on warning and shaking they rapidly become crystalline. The salts thus obtained are hydrated, I molecule of the oxalate being as a rule associated with 9, 10, or 11 molecules of water.

The ovalates of lanthamm, cerum, and "didymmm" of the type  $M_2(C_2O_4)_3$  11H<sub>2</sub>O are isomorphous, crystallising in the monochine system:—

|                                           | а      | b | •      | β            |     |
|-------------------------------------------|--------|---|--------|--------------|-----|
| $\text{La}_{2}(C_{2}O_{4})_{3}.1111_{2}O$ | 1 0730 | 1 | 21750  | $92^{\circ}$ | 27  |
| Ce.j(C.O.) 1111.jO                        | 1 1300 | ı | 20098  | 95           | 13' |
| "Để (C.Ő.), 11H.O                         | 1 0693 | 1 | 2.1346 | 92           | 17  |

and the enneabydrates of lanthanum and cernm,  $M_2(C_2O_1)$ ,  $911_2O_1$  are also isomorphous, crystallising in the tetragonal system 3

The oxalates are practically insoluble in water The following table contains the solubility determinations that have been published, the data representing milligrams of anhydrous ovalate per litre at 19° 27° C = -

| Solid Phase                                                                                                                                                                                                                                                                                                                               | ļ | Λ                            | В            | C            | D.           |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---|------------------------------|--------------|--------------|--------------|
| La <sub>2</sub> (C <sub>2</sub> O <sub>4</sub> ), 10H,O .<br>Ce <sub>1</sub> C <sub>2</sub> O <sub>4</sub> 10H,O                                                                                                                                                                                                                          | • | 0 62<br>0 11                 | 0 70<br>0 15 | 0 96<br>0 81 | 1 21<br>0 99 |
| P <sub>1</sub> (C <sub>2</sub> O <sub>4</sub> ) <sub>1</sub> 10H <sub>2</sub> O<br>Nd <sub>2</sub> (C <sub>2</sub> O <sub>4</sub> ) <sub>1</sub> 10H <sub>2</sub> O<br>Sm <sub>2</sub> (C <sub>2</sub> O <sub>4</sub> ) <sub>2</sub> 10H <sub>2</sub> O<br>Y <sub>2</sub> (C <sub>2</sub> O <sub>4</sub> ) <sub>1</sub> 9H <sub>2</sub> O |   | 0.71<br>0.49<br>0.51<br>1.00 | 0 31         | 0.59         | 0.73         |
| "Yb2"(C2O4 10H3O                                                                                                                                                                                                                                                                                                                          |   | 3 34                         |              |              |              |

The values given under A were determined by Rimbach and Schubert 1 by the electrical conductivity method. The other values are due to Hauser and Herzfeld , those given under B were determined from conductivity measurements, and those under C and D by the gravimetric and volumetric analyses of the saturated solutions. The determinations are only approximately correct, but they show the ovalates of the yttrium group to be more soluble in water than those of the cerum group

The oxalates are slightly soluble in dilute immeral acids, the solubilities increasing with the concentration of the acid and with the temperature. The solubilities in aqueous sulphuric acid at 25° have been carefully determined for a few oxalates, 6 the results are shown graphically in fig. 21, in which, \_\_\_\_\_

- -

<sup>1</sup> On the precipitation of the pure ovalates, see the section on analytical chemistry

<sup>1</sup> On the precipitation of the pure oxalates, see the section on analytical chamistry (p. 368)
2 Wyrouboff, Bull Soc franç. Min., 1901, 24, 105, 1902, 25, 66.
3 Wyrouboff, thad, 1 101, 24, 111
4 Rimbach and Schubert, Zeitsch physikal Chem., 1909, 67, 183
5 Cited in Meyer and Hauser, Die Analyse der seitenen Erden und der Erdsauren (Stuttgart, 1912), p. 61
6 Hauser and With, Zeitsch anal. Chem., 1904, 47, 389, Wirth, Zeitsch anang Chem., 1912, 76, 174; Meyer and Wassjuchnow, cited in Meyer and Hauser, opus cit. (Nd.), cf. Brainer, Trans. Chem. Soc., 1898, 73, 951.

instead of plotting the amounts of the oxalates contained in the solutions, the equivalent qualities of the oxides have been indicated. The solubilities of scandium and thorium oxalates have also been indicated for purposes of comparison. Were it not for the position of the erbium oxidate curve, it

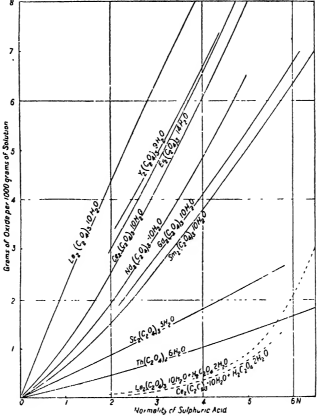


Fig. 21.—Solubilities of the exalates of some of the rate earth elements in aqueous sulphune acid at 25° C.

might be concluded that the more basic the rare earth, the more soluble is

the corresponding oxalate in sulphuric acid.

A few of the experimental results for the solubilities are contained in the following table, the data denoting grams of oxide (CeO, in the case of cerium) or anhydrous oxalate per 100 grams of saturated solution:-

## THE RARE EARTH ELEMENTS.

| Normality      |                                   | y of Oxalate<br>erms of                           | Normality<br>of   | Solubility of Oxalate<br>in Terms of |                       |  |
|----------------|-----------------------------------|---------------------------------------------------|-------------------|--------------------------------------|-----------------------|--|
| Sulphunc Acid  | Oxide                             | Auhydrous<br>Oxalate,                             | Sulphune<br>Acid. | Oxide                                | Anhydrous<br>Oxalate, |  |
|                | La <sub>2</sub> (C <sub>2</sub> C | ,), 10H <sub>2</sub> O                            |                   | Sm <sub>2</sub> (C <sub>2</sub> O    | 7 10H2O               |  |
| 0 10           | 0.0208                            | 0 0346                                            | 0 10 i            | 0.0058                               | 0.00901               |  |
| 0.50           | 0 0979                            | 0.1628                                            | 0.50              | 0 0313                               | 0 0507                |  |
| 1 00           | 0 2383                            | 0 3962                                            | 1 00              | 0 0627                               | 0 1016                |  |
| 1 50           | 0.3190                            | 0 5304                                            | 1.415             | 0.1114                               | 0 1804                |  |
| 2 00           | 0 1417                            | 0 7344                                            | 2 39              | 0 1914                               | 0 3099                |  |
| 3 20           | 0 7632                            | 1 2690                                            | 4 32              | 0 4328                               | 0 7008                |  |
|                | Ce2(C,C                           | ), 10H <sub>2</sub> O.                            |                   | Gd.(('_()                            | ا0ر1101 بن            |  |
| 0.10           | 0 0136                            | 0.0215                                            | 2 16              | 0 1883                               | 0 3005                |  |
| 0.50           | 0 0524                            | 0 0828                                            | 311               | 0 3010                               | 0 4503                |  |
| 1 00           | 0 1140                            | 0 1802                                            | 4 32              | 0 1359                               | 0 6956                |  |
| 1 ·445<br>2 39 | 0 1764<br>0 3083                  | 0 2788<br>0 4873                                  |                   | O,''),.Y                             | , oli ,O.             |  |
| 3 90           | 0 6300                            | 0 9957                                            | 2 16              | 0.352                                | 0 6584                |  |
| 3 30           |                                   | ) <sub>4</sub> ) <sub>3</sub> 10H <sub>2</sub> O. | 1 32              | 0 7236                               | 1.1000                |  |
| 0.50           | 0 0336                            | 0 0552                                            |                   | Er2(('2()                            | 0رااا1 ا              |  |
| 1 00           | 0 0752                            | 0 1235                                            | 2 16              | 0:329                                | 0.514                 |  |
| 2 00           | 0 1872                            | 0 3071                                            | 3 11              | 0 193                                | 0 771                 |  |
| 5.00           | 0 6603                            | 1 0840                                            | 1 32              | 0 704                                | 1 100                 |  |

When the concentration of the sulphinic acid exceeds a certain value, the stable solid phase in contact with the solution changes from the oxalate to the sulphate 1

The Syalates of the care each elements are more soluble in aqueous hydrochloric than in sulphuric acid, differing in this respect from scandium oxalate (see p 214) This will be seen from fig 22, where the results for cerous, needymnum, and scandium oxalates are represented graphically. The numerical data 2 are as follows, in grains of authydrous oxalate per 100 grams of solution -

| Normality of HCl                                                                                                                                                     | 0.125. | 0 25   | 05               | 10     | 1.5   | 2              | 3     | 4     | 5     |   |
|----------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------|--------|------------------|--------|-------|----------------|-------|-------|-------|---|
|                                                                                                                                                                      | · '    |        |                  |        | -     |                |       | !     |       | l |
| Ce <sub>2</sub> (C <sub>2</sub> O <sub>4</sub> ) <sub>3</sub> 10H <sub>2</sub> O<br>Nd <sub>2</sub> (C <sub>2</sub> O <sub>4</sub> ) <sub>3</sub> 10H <sub>2</sub> O | 0 0151 | 0.0313 | 0 0812<br>0 0270 | 0.1971 | 0 355 | 0 532<br>0 333 | 0 977 | 1 555 | 1 721 |   |

From hot, concentrated hydrochloric acid, the oxalochlorides of the rare

earth elements,  $M(C_2O_1)C(\mathbb{R}^{3}H_2O)$ , may be obtained <sup>1</sup> The oxalates of the rare earth elements are practically insoluble in aqueous oxalic acid, thereby resembling thorium and scandium oxalates, but differing from the zirconium salt. Moreover, the addition of a moderate amount of ovalic acid greatly reduces the solubilities of the ovalities in mineral acids, a fact of great practical importance. The extent by which the

See Wiith, Zeitsch anorg. Chem., 1908, 58, 213.
 Taken from Meyer and Hauser, opus cit.
 Job, Compt. rend., 1898, 126, 246.

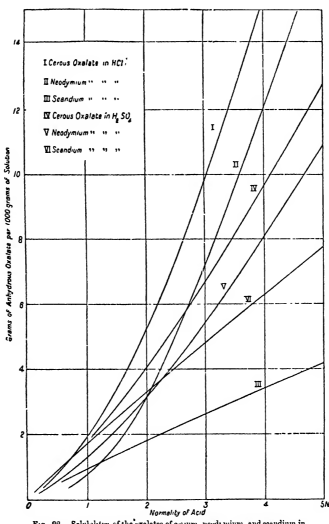


Fig. 22.—Solubilities of the exalates of cerum, needymium, and scandium in aqueous hydrochlone and sulphuric acids at 25° C.

solubility is diminished may be seen by comparing the results given in the following table  $^1$  with those already quoted (solubilities in grains of anhydrous exalate per 100 grains of solution at  $25^{\circ}$  ( $^{\circ}$ ) —

| Norm                                                                          | ality of | Solubility of                                                        |                    |                                                                      |  |  |  |  |
|-------------------------------------------------------------------------------|----------|----------------------------------------------------------------------|--------------------|----------------------------------------------------------------------|--|--|--|--|
| H <sub>2</sub> SO <sub>4</sub>   H <sub>2</sub> C <sub>2</sub> O <sub>4</sub> |          | L <sub>12</sub> (C <sub>2</sub> O <sub>4</sub> ), 10H <sub>2</sub> O | Ce2(('2O4), 10112O | Sm <sub>2</sub> (C <sub>2</sub> O <sub>4</sub> ), 10H <sub>9</sub> O |  |  |  |  |
|                                                                               | -        | <u></u>                                                              | ; —                | ¦                                                                    |  |  |  |  |
| 0 05                                                                          | 0 05     |                                                                      | 0 0030             | [                                                                    |  |  |  |  |
| 0 50                                                                          | 0 05     | 0 0334                                                               | 0 0105             |                                                                      |  |  |  |  |
| 0 25                                                                          | 0 25     | 0 0099                                                               | 0 0016             |                                                                      |  |  |  |  |
| 0 05                                                                          | 0.50     | 0 0039                                                               | 0.0025 2           | 0.0009                                                               |  |  |  |  |
| 0 25                                                                          | 0.50     | 0 0069                                                               |                    |                                                                      |  |  |  |  |
| 0 50                                                                          | 0.50     | 0 0090                                                               | 0.0010 2           | 0 0010                                                               |  |  |  |  |
| 0 96                                                                          | 0.50     | 0 0138                                                               | 1                  | 0 0032                                                               |  |  |  |  |
| 1.19                                                                          | 0.50     | 0 0203                                                               |                    | 0 0042                                                               |  |  |  |  |
| 1.44                                                                          | 0.50     | 0.0293                                                               | ·                  | 0 0057                                                               |  |  |  |  |
| 1 70                                                                          | 0.50     | 0 0369                                                               | ł .                | 0 0080                                                               |  |  |  |  |

The diminution in the solubility of fanthanum and cerous ovalates in aqueous sulphunic acid, brought about by saturating the solutions with ovale neid, is shown graphically in fig. 21 by the dotted lines, which represent the results of Wirth

The evalutes of the cerum group are practically insoluble in dilute aqueous solutions of alkali evalutes, those of the yttimm group are perceptibly soluble. One grain of ammonium evalute dissolved in 38 grains of water at 20° C. dissolves the evalutes in quantities equivalent to the following amounts of evides  $-\frac{\pi}{2}$ 

| Oxide                                    | [m,O3.  | P12O1          | Nd_O <sub>3</sub> | ('e <sub>2</sub> () | Y <sub>2</sub> O <sub>4</sub> | "O"cdY          | ThO <sub>2</sub> |  |
|------------------------------------------|---------|----------------|-------------------|---------------------|-------------------------------|-----------------|------------------|--|
| Grams dissolved<br>Relative solubilities | 0 00023 | 0 00023<br>1 1 | 0 00034           | 0 00012             | 0 00256<br>11 0               | 0 0214<br>101 6 | 0 620<br>2663    |  |

It is clear that the order of increasing solubility of the oxidates is approximately the order of decreasing basic strength of the corresponding oxides. The slight solubilities of the oxidates of the rare earth elements, in comparison with the solubility of thorium oxidate, is also obvious

Ammonium and potassium oxalates are better solvents for the oxalates of the yttrium group than sodium oxalate. The solubility of the oxalates of the irre earth elements in solutions of these salts is connected with the fact that double or complex oxalates are produced. In the yttrium group the double ammonium oxalates may be readily prepared by dissolving the rare earth oxalates in hot, concentrated ammonium oxalate and cooling the solutions, the salts thus obtained are of considerable value for the separation of the yttria earths. Various double oxalates are stated to be formed merely

<sup>1</sup> Hauser and Wirth, loc, cit

These results appear to be in error Brauner, Trans Chem Soc, 1898, 73, 951.

by the addition of soluble salts of the rare earth elements to concentrated solutions of ammonium or an alkali oxalate (see pp. 425, 431).

In consequence of this tendency towards the formation of double oxalates, occlusion of soluble oxalate occurs to a notable extent when the oxalates of the rare earth elements are precipitated by means of soluble exalates instead of oxalic acid. This is the case even with the oxalates of the cerium group.1

It has been mentioned that oralorhlorides may be prepared by crystallising solutions of the oxalates in hot, concentrated hydrochloric acid. These salts may also be prepared by dissolving the oxalates in concentrated solutions of the corresponding chlorides,2 Analogous oxalobromides and oxalo-iodides are stated by Job to exist The ovalates are about as soluble in dilute nitric as in dilute hydrochloric acid, and may be crystallised unchanged from the solution, but when heated with concentrated mitric acid decomposition ensues and oxalonitrates are produced.3 On further heating with boiling, concentrated nitric acid, the oxalonitrates are readily oxidised to nitrates. The conversion of oxalate into nitrate in this manner proceeds very readily in the case of the cerium salt, the presence of which also has a very marked catalytic effect in increasing the rates of oxidation of the other oxalates 4 Oxalonitrates may in certain cases be produced by dissolving the oxalates in concontrated solutions of the corresponding intrates (sec, eg., p. 420). It is probable that oxulosulphates may also be prepared.

When heated, the oxalates first lose their water of crystallisation, but it is almost impossible to obtain the pure, anhydrous oxalates, decomposition commencing before the last traces of water are expelled. In decomposing, the oxalates first evolve carbon monoxide, leaving behind the carbonates, and the latter then lose carbon dioxide and leave the oxides. When thus heated to redness in air, cerous oxalate leaves a residue of ceria, CeO, praseodymum and terbium oxalates leave peroxides of the (approximate) composition M,O, and the other oxalates leave the sesqui-oxides, M,O, as

The oxalates of the rare earth elements are extremely important substances from the practical point of view. At times they may be conveniently propared by heating the precipitated hydroxides, carbonates, or double alkali sulphates with aqueous ovalie acid. It is often necessary to transform the oxalates into soluble salts. The conversion into mitrates by heating with nitric acid has been mentioned. The oxalates may be transformed into anhydrous sulphates by heating with concentrated sulphutic acid. By heating with concentrated alkali hydroxide (preferably potassium hydroxide) they may be converted into the hydroxides, which may then be dissolved in a suitable acid. The transformation from oxalate to another salt may often be readily carried out by first igniting the oxalate to oxide and then dissolving the latter in an acid, unless a mixture of oxides is obtained containing more than 45-50 per cent of certa!

Ethylsulphates, M(C, H5, SO4), 9H2O - The ethylsulphates of the rare earth metals are beautifully crystalline salts They may be prepared by two

<sup>1</sup> Baxter and Guiffin, J. Amer. Chem Soc., 1906, 28, 1684, Baxter and Daudt, abid.,

<sup>1908, 30, 563.</sup> See also p. 368

Matignon, Ann. Chim. Phys., 1906, [viii ], 8, 243.

Biauner, Trans. Chem. Soc., 1898, 73, 951; R. J. Meyer and Marckwald, Ber.,

<sup>1900, 33, 3008.</sup>Baibieri and Volpino, Attr R. Accad Lincer, 1907 [v.], 16, i. 399. <sup>5</sup> For the necessary procedure in such a case, see p. 332.

methods: (i) double decomposition between barium ethylsulphate and the rare earth sulphates, rigorously exempt from free acid, and crystallisation of the liquid at as low a temperature as possible after removing the barnim sulphate, 1 and (11) double decomposition between alcoholic solutions of sodium ethylsulphate and the rare earth chlorides, sodium chloride being precipitated 2

The ethylsulphates form a strictly isomorphous series of salts, crystallising from water as the hydrated salts M(C<sub>2</sub>H<sub>5</sub>SO<sub>1</sub>)<sub>3</sub>,9H<sub>2</sub>O, which belong to the bipyramidal class of the hexagonal system. Twelve of them have been carefully examined by Jaeger, to whom the following data are due -

| Salt of | Density<br>at 25° | Molec.<br>Volume. | Axial Ratios a c. | Salt of Done |          | Axial Ratios |
|---------|-------------------|-------------------|-------------------|--------------|----------|--------------|
| l       |                   |                   |                   |              |          |              |
| Li      | 1 845             | 366.6             | 1 0 5073          | Gd4 : 1:91   | 19 362 0 | 1:0 5050     |
| Ce      | 1 930             | 351 1             | 1 0 5075          | Dy 194       | 2 360 4  | 1 0.5050     |
| Pr      | 1.876             | 361:1             | 1 0 5059          | Y 1.70       | 354 9    | 1:0 5035     |
| N43     | 1 883             | 362 1             | 1 0 5068          | Fr 190       | 369.8    | 1 0 5053     |
| Sur     | 1 904             | 361 3             | 1 0 5072          | Tm 2 00      | 01 352 8 | 1 0 5014     |
| Ea      | 1 909             | 361 2             | 1 0 5058          | Yb 2 01      | 19 351 4 | 1 0 5079     |
| 1       | i                 | 1                 |                   |              |          |              |

According to Jacger, in no case does the axial ratio  $a \cdot c$  differ from the mean value 1  $0.5062 \pm 0.0012$  by more than can be attributed to experimental error Scandium ethylsulphate, moreover, is not isomorphous with the preceding salts 5

The ethylsulphates are extremely useful salts to use for the fractional crystalliation of the earths of the yttimm group

Acetylacetonates, MCH CO CH CO CH<sub>1</sub>)<sub>3</sub> --These substances may be prepared by shaking the hydroxides with a dilute alcoholic solution of acetylactione, or, more simply, by adding a slightly ammoniacal solution of acetylactione to a neutral solution of the chloride or intrate. They are sparingly soluble in water, but may be crystallised from aqueous alcohol or organie solvents such as benzene, chloroform, etc

The melting points of the acetylacetonates are not very definite. W. Biltz gives the following values ---

The cerous compound forms a trihydrate, m p. 145° C  $^{\circ}$ The acetylacetonates cannot be sublimed without decomposition taking place. In this respect, and

Urham, Bull Sec, chim., 1898, [m], 19, 378., Ann. Chim. Phys., 1900, [vn.], 19, 184., J. Chim. phys., 1906, 4, 56
 James, J. Amer. Chem. Soc., 1912, 34, 767
 Cf. Morton, Zulsch. Kryst. Min., 1887, 12, 517.
 Cf. Benedicks, Zulsch. anney Chem., 1900, 22, 413
 Jaeger, Proc. K. Akad. Weienwh. Amsterdam., 1914, 16, 1095. Rec. trav. chim., 1914, 2006.

<sup>33, 342;</sup> July Soc chim., 1897, [iii.], 17, 98; Ann Chim Phys, 1909, [vii.], 19, 184; Ulbain and Buchschovsky, Compt. rend., 1897, 124, 618; Handwich and Desch, Annalen, 1902, 232; 1 (La), W. Biltz, ibid., 1904, 331, 344 (La, Cc, Pr, Nd, Sm), W. Biltz and Chinch, Zeitsch anorg Chem., 1904, 40, 218, James, J. Amer. Chem. Soc., 1911, 33, 1382 (Tm); G. T. Morgan and H. W. Moss, Trans. Chem. Soc., 1914, 105, 189 (Y).

in crystalline form, they differ from the scandium compound. In solution they exist mainly as double molecules, M2(CH, CO CH CO CH,), The acetylacetonates form crystalline, addition compounds with aminoma, substituted ammomas, and pyridine

Other Organic Salts. In the hope of finding sintable compounds for effecting the fractionation of the rare earths, numerous organic salts of rare earth elements have been prepared 3

The malonates are sparingly soluble crystalline salts, prepared from the hydroxides of the metal and malonic acid. The following have been

```
\begin{array}{c} \text{La}_2[\text{CH}_2(\text{COO})_2] \ 5\text{H}_2\text{O} \\ \text{Ce}_2[\text{CH}_2(\text{COO})_2] \ 5\text{H}_2\text{O} \\ \text{Pr}_2[\text{CH}_2(\text{COO})_2] \ 6\text{H}_2\text{O} \\ \text{Nd}_2[\text{CH}_2(\text{COO})_2] \ 5\text{H}_2\text{O} \end{array}
   \begin{array}{l} {\rm Sn_{*}[CH_{2}(COO),].6H_{2}O} \\ {\rm Gd_{2}[CH_{2}(COO),].8H_{2}O} \\ {\rm Y_{2}[CH_{2}(COO),].5H_{2}O} \\ {\rm Er_{2}[CH_{2}(COO),].10H_{2}O} \end{array}
```

In the ceruin group the salts form monochnic plates, in the yttirum group they crystallise in thombic needles 1

The glycollates, like the malonates, are sparingly soluble crystalline salts. In the cerum group the salts are anhydrons. The following solubilities, in grams of salt per litre of solution at 18', are due to Jantsch

```
La[CH_cOH)COO]<sub>3</sub> 3 328
   \begin{array}{ccc} {\rm Sm}[{\rm CH}_2({\rm OH})\;{\rm COO}] & -6\;373 \\ {\rm Gd}[{\rm CH}_2({\rm OH})\;{\rm COO}]\; 2{\rm H}_2{\rm O} & -11\;117 \\ {\rm Y}[{\rm CH}_2({\rm OH})\;{\rm COO}]\; 2{\rm H}_2{\rm O} & -2\;417 \end{array}
```

The cacodylates are of interest, since those of the cermin group are readily soluble and those of the yttrum group practically insoluble in cold water The solubilities at 25, in grams of salt dissolved by 100 grams of

```
\begin{array}{lll} \operatorname{La}[(\operatorname{CH}_4)_2\operatorname{AsO}_2]_4\,x\operatorname{H}_2\operatorname{O} & \operatorname{v. sol} \\ \operatorname{Ce}[(\operatorname{CH}_4)_2\operatorname{AsO}_2]_4\,x\operatorname{H}_2\operatorname{O} & \operatorname{v. sol} \\ \operatorname{Pr}[(\operatorname{CH}_4)_2\operatorname{AsO}_2]_4\,8\operatorname{H}_2\operatorname{O} & 8 \ 13 \\ \operatorname{Nd}[(\operatorname{CH}_4)_2\operatorname{AsO}_2]_3\,8\operatorname{H}_2\operatorname{O} & 5 \ 13 \end{array},
   \begin{array}{lll} \mathrm{Sm}[(\mathrm{GH}_3)_2\mathrm{AsO},] \; 8\mathrm{H}_3\mathrm{O} & 1 \; 60 \\ \mathrm{Y}[(\mathrm{GH}_4)_2\mathrm{AsO}_2] \; 9\mathrm{H}_3\mathrm{O} & \mathrm{insol} \\ \mathrm{Tm}[(\mathrm{GH}_3)_2\mathrm{AsO}_2] \; 8\mathrm{H}_2\mathrm{O} & \mathrm{insol} \end{array}
```

The cacodylates form double salts with the rare earth chlorides, intrates, etc., they are crystalline and sparingly soluble in water. The lanthanum, cerum, and neodymium salts of the type 2M[(CH<sub>3</sub>),AsO<sub>2</sub>], MCl<sub>3</sub>18H<sub>2</sub>O have been

<sup>1</sup> Jacget, loc cit.
2 See p 236
3 A list will be found under the heading of each of the rate earth elements, see Chaps.
XII., XIII., and XIV
4 Endmann and Wuth, Annalen, 1908, 361, 190 (La. Ce. Pl. Nd. Sni Gd. Y. Er), Rimbach and Kilian, ibid., 1909, 368, 110 (Ce), Holmberg. Zeitsch annag Chem., 1907, 53, 29 (La. Ce. V).

Rimbach and Kilian, 1014, 1903, 306, 110 (Co), Assessing the Second State of Rimbach and Giunkiaut, Zeitsch anorg, Chem., 1912, 79, 305. Second Rimbach and Kilian, Annalen, 1903, 368, 110 (Ce), James, Hoben, and Robinson, J. Amer. Chem. Soc., 1912, 34, 276 (Sm), Pratt and James, thid., 1911, 33, 13:0 (Y).

8 Whittemore and James, J. Amer. Chem. Soc., 1913, 35, 127.

The sebacates are of interest owing to the fact that they are of value in quantitative analysis — The following have been described —  $^1$ 

$$\begin{array}{lll} \text{La}_{2}[C_{3}\Pi_{10}(\text{COO})_{2}]_{1}\,2\Pi_{10} & \text{Sm}_{2}[C_{3}\Pi_{10}(\text{COO})_{2}]_{1}\,4\Pi_{2}O \\ \text{Pr}_{2}[C_{3}\Pi_{10}(\text{COO})_{2}]_{2}\,2\Pi_{10} & \text{Y}_{2}[C_{3}\Pi_{10}(\text{COO})_{2}]_{3}\,4\Pi_{2}O \\ \text{Nd}_{2}[C_{3}\Pi_{10}(\text{COO})_{2}]_{3}\,3\Pi_{2}O \end{array}$$

The dimethylphosphates, M[(CH), PO<sub>1</sub>], are of considerable practical uc. In the cerum group they are readily soluble and in the yttrium group sparingly soluble in water, and all the salts are less soluble in hot water than in cold With the exception of the lanthanum and cerium salts, they are anhydrous In the cerium group the salts crystallise in hexagonal plates or prisms, but in the terbium and yttrium groups they crystalliso in long needles. The solubilities, in grams of anhydrous salt per 100 grams of water at 25° and 95°, are as follows =2

| Salt of        | 25°                   | 95° | Salt of        | 25*                  | 95                  | Silt of            | 25°                | 95°.         |
|----------------|-----------------------|-----|----------------|----------------------|---------------------|--------------------|--------------------|--------------|
| La<br>Ce<br>Pi | 103 7<br>79 6<br>64 1 | 65  | Nd<br>Sm<br>Gd | 56 1<br>35 2<br>23 0 | 22 3<br>10 8<br>6 7 | <br>Y<br>Et<br>Y b | 2 8<br>1 78<br>1 2 | 0 5!<br>0 25 |

The m-nitrobenzenesulphonates,  $M[C_0H_1(NO_2)SO_1]_p$  are of value for the separation of the cerum group. They are reachly soluble in water and crystallise readily. Two isomorphous series are known. (i.) the hexahidrates of the La, Ce, Pi, and Nd salts, and (ii.) the heptahidrates of the Sin, Gd\_and Y salts. The solublities at 15, in grains of anhydrous salt per  $M_1(NO_1)$  and  $M_2(NO_2)$  are of value. 100 grams of water, are as follows - 1

The  ${\tt I}$ :4:2-bromontrobenzenesulphonates are well-defined crystilline salts, crystallising with 8, 10, or 121120. The following have been prepared --

The solubilities at 25° (expressed as grains of anhydrous salt in 100 grains of saturated solution) are as follows — "

| La salt | 177  | Sm salt | 7 27 | Er salt | 6 06 |
|---------|------|---------|------|---------|------|
| Ce "    | 5.56 | Eu "    | 6.31 | Tm ,,   | 6.38 |
| Pr "    | 5 73 | Gd "    | 5 94 | Yb "    | 729  |
| N' J "  | c 7c | v       | 5.74 |         |      |

The p-dichlorobenzenesulphonates and p-dibromobenzenesulphonates of a number of rare earth elements have been examined

Whittemore and James, loc cit
 J. C. Morgan and James, J. Amer. Chem. Soc., 1914, 36, 10
 Holmberg, Zeitsch anony Chem., 1997, 53, 83
 Katz and James, J. Amer. Chem. Soc., 1914, 35, 872
 See also fig. 35 on p. 326.

crystallographically by Armstrong and Redd.<sup>1</sup> The dibromo salts of the cerium group form hydrates with 9H<sub>2</sub>O and with 18H<sub>2</sub>O (Sm salt not known). The enneahydrates form a series of isomorphous, orthorhombic salts :-

```
a:b c.
\begin{array}{c} {\rm La}({\rm C_6H_3Fr_2.SO_8)_3~9H_2O} \\ {\rm Ce}({\rm C_6H_3Br_2~SO_3)_3.9H_2O} \\ {\rm Pr}({\rm C_6H_3Br_2~SO_3)_3~9H_2O} \end{array}
   1.3965 - 1 - 0 8753
  14106 1 08873
   1:3964:1:08798
Nd(C_6H_3Br_2.SO_3)_3 9H_2O
  1.3990 1.08789
```

Gadolinium, however, forms a monoclinic heptahydrate,  $Gd(C_6H_3Br_aSO_3)_3$   $7H_3O$  (a:b c=1.2595.1 06031;  $\beta$ =89° 16′) and a monoclinic dodecahydrate,  $Gd(C_6H_3Br_2SO_3)_3$   $12H_2O$ . The latter is isomorphous with the praseodyminin and neodymium salts of p-dichlorobenzene sulphonic acid -

|                                                                                                    | abr                       | β       |
|----------------------------------------------------------------------------------------------------|---------------------------|---------|
| Pr(C <sub>6</sub> H <sub>8</sub> Cl <sub>2</sub> SO <sub>3</sub> ) <sub>3</sub> 12H <sub>2</sub> O | $0.5887 \cdot 1 : 0.3819$ | 76° 26′ |
| $Nd(C_6H_3C_2SO_3)_812H_9O$                                                                        | 0.5872 1 · 0 · 3810;      | 76° 34′ |
| Gd(C,H,Br.SO,), 12H,O                                                                              | 0.5952:1 0.3817.          | 76° 18′ |

Further, lanthanum and prascodymium form isomorphous, triclinic pentadecahydrates, La(U<sub>0</sub>H<sub>1</sub>,U<sub>2</sub> SO<sub>1</sub>)<sub>3</sub>, 15H<sub>2</sub>O (a b, c=1 6193 1 1 6028, a = 76° 26′,  $\beta=113^\circ$  48′,  $\gamma=68^\circ$  6′) and Pr(U<sub>0</sub>H<sub>3</sub>Cl<sub>2</sub> SO<sub>2</sub>)<sub>3</sub> 15H<sub>2</sub>O

Various tartrates,2 citrates,2 malates,8 etc., have also been examined

# THE RARE EARTH ELEMENTS AND BORON

Metaborates, M(BO<sub>2</sub>)<sub>3</sub>—When the oxalate of a rare earth element is added to excess of molten boron sesqui-oxide, the mass obtained separates into two layers (conjugate phases), the upper one consisting almost entirely of boron sesqui-oxide. The lower layer solidifies to a glassy mass, which devitrifies when heated for some hours over a Bunsen burner. From it a little bone anhydride can then be extracted with boiling water, leaving as residue a crystallino metaborate,  $M^{11}(BO_2)_3$ . Even in the case of cerum, the product is said to be of this type, if the cerous oxalate is added to the molten oxide in an atmosphere of carbon dioxide, and it is also stated that cerous metaborate is obtained when ceric oxide is substituted for cerous oxalate 4

## SPECTRA OF THE RARE EARTH ELEMENTS AND THEIR COMPOUNDS 5

The various types of rare earth spectra may be classified as follows -

- A. Absorption spectra.—These include the following
  - (a) Absorption spectra by transmission.
  - (b) Absorption spectra by reflection.

<sup>&</sup>lt;sup>1</sup> H. E. Armstrong and Rold, Proc. Roy. Soc., 1912, A, 87, 204, Rodd, abid, 1914, A,

<sup>1</sup> H. E. Armstrong and Rodd, Proc. Roy. Soc., 1912, 21, 21, 22, 23, 292.
2 Holmberg, loc. cit.
3 Jantsch, Chem. Zeit, 1914, 38, 794
4 Guertler, Zeitsch anorg Chem., 1904; 40, 225 (La, Ce, ''D, ''Sm, Gd), cf. Nordenskield, Poyy. Annalen, 1861, 114, 612, Holm, Inaugural Dissettation (Munich, 1902)
5 The most comprehensive treatise on Spectroscopy is Kayes, Handbuch der Spektroskopis (Hirzel, Leipzig, 6 vols., 1900-1910). Of the smaller text books, Baly, Spectroscopy (Longmans, 2nd ed., 1912), and Urbain, Introduction à l'étude de la spectrochimic (Heimann et Fils, Paris, 1911), may be mentioned, the latter gives a very clear account of the theoretical side of the subject.

### B. Emission spectra.—These include the following .-

- (1.) Bund spectra.
  - (a) Flame spectra
  - · (b) Reversion spectra (spectres de renrer sement).
  - (c) Cathodic phosphorescence spectra.
- (ii ) Line spectra
  - (a) Spark spectra
  - (b) Are spectra
  - (c) X-ray or high frequency spectra

The above order of arrangement is adopted in the following account.

## ABSORPTION SPECTRA BY TRANSMISSION AND REFLECTION

General.—The salts (derived from colourless acids) of the following rare earth elements are coloured, and in solution exhibit well-defined absorption spectra in the visible region praseodymium, neodymium, samarium, europium, dysprosium, holmium, erbium, and thulium. In addition, the salts of terbium exhibit a single band in the visible region of the spectrum. Absorption bands are not limited merely to the visible part of the spectrum, but may also occur in the infra-red and ultra-violet regions. The absorption spectrum of gadolinium, for instance, lies wholly in the ultra-violet

The absorption spectra of the rare earth compounds differ markedly from other absorption spectra in the large number of bands they exhibit, and in the great intensity and the remarkable narrowness of many of the bands. Of the visible absorption spectra, only those of europium and terbium are weak. The presence of the absorbing rare earth elements is therefore easily detected. Further, when the absorption spectrum of a rare couth element appears only family, the element is known to be present only in very small concentration

The literature relating to the absorption spectra of the compounds of the rate earth elements is fairly extensive. It may be recalled that the first observations on these spectra were made by Bahr in 1852 and that their study was pursued by Gladstone, Delafontaine, and others. At a period when the discovery of samarskite had led to renewed interest in the rare earths, the work of Soret upon their absorption spectra was of great importance 1. From the historical point of view, the work of Kruss and Nilson 2 is also of interest Of the more modern work, that of Forsling, and particularly that of H. C. Jones and his co workers,4 may be mentioned

The absorption spectra of solids may be examined in the ordinary manner applicable to liquids if thin, transparent, crystal slices are available, other

Soret, Compt. rend., 1878, 86, 1062, 1879, 88, 422, 1077, 89, 521, 1880, 91, 378;
 Arch Scz. phys. nat., 1878, [ii.], 63, 11, 89, 1880, [iii.], 4, 261.
 See p. 287.
 Forsling, Bilwing K. Szenska Vet.-Akad Handl., 1892, 18, I., No. 4, 1893, 18, I., No. 10, 1892, 23, I., No. 5, 1899, 24, I., No. 7, 1902, 28, II., No. 1
 Jones and Anderson, Proc. Amer. Phil. Soc., 1998, 47, 276, Amer. Chem. J., 1909, 41, 276, Jones and Strong, Proc. Amer. Phil. Soc., 1909, 48, 194, Amer. Chem. J., 1910, 43, 37, 97, 1911, 45, 1, 1912, 47, 27, 120, Jones and Guy, ibid., 1913, 49, 1, 50, 257; Physikal Zetterh. 1912, 13, 619, Jones and Anderson, Carnegie Institution Publications, 1909, No. 110, Jones and Strong, ibid., 1915, No. 210

wise it is necessary to examine the light they reflect for absorption bands. Observations upon the absorption spectra of the rare earths are usually made with salt solutions, generally with aqueous solutions

Absorption spectra are often described by stating for each band the limits between which absorption is observed in the spectrum. This, however, necessitates a full description of the experimental details to be of real value, since the width of a band is increased when either the concentration of the solution or the depth of solution through which absorption takes place is increased. Moreover, the edges of the bands are often hazy and difficult to define with accuracy. A better method of description is to state for each band the wave length corresponding to the position of maximum absorption, i.e. what is known as the "head" of the band. This is not necessarily at the middle of the band, moreover, a broad band may show two or more maxima. The position of the "head" of a band is, in general, practically independent of the concentration of the solution.

The absorption is said to follow *Beer's Law* when the increase in the intensity of absorption at any point in the spectrum due to increase of concentration of the absorbing solution, may be exactly millihed by a diministron in thickness of the absorbing solution in the same ratio. For incdentely dilute solutions (aqueous or otherwise) of race earth compounds, the absorption is described with considerable accuracy by Beer's Law, as If C Jones and his co-workers have shown

When the absorption spectrum of a rare earth compound in solution is observed at different dilutions through the same thickness of solution, it is observed that certain broad bands seen in concentrated solution become resolved into a number of narrower bands as the dilution is continued. Eventually, as dilution proceeds, the bands disappears but the dilution at which one band disappears is not necessarily the same as that at which any other band ceases to be visible. Some bands can only be observed in moderately concentrated solution, while others are still visible in extremely dilute solution. It must therefore be borne in mind when consulting complete lists of absorption bands that all the bands eminicated cannot be seen at one and the same time, for in solutions sufficiently concentrated to exhibit the weak bands others will have coalesced, producing a similler number of broader bands (see figs. 24 and 25 on pp. 288 and 290).

When the absorption spectra of various compounds of the same rare earth element are examined in the absence of any solvent, it is found that they are decidedly different from one another, although a certain "family likeness" may be apparent. It was discovered by Binsen that the relative intensities of the bands in the spectrum of "didynum" sulphate octahydrate vary according to direction in which light passes through the crystal. This arises from the fact that the crystal is doubly refracting and absorbs the ordinary and extraordinary rays unequally, i.e. the crystal is pleechine. If Becquerel has studied this phenomenon in great detail for various salts of "didynum," and also for various transparent innerals, such as scheelite, apartite, parisite, xenotime, monizate, etc., which contain "didynium" either in traces or in quantity. The bands differ only in relative intensities, not in actual position.

Bunson, Popt. Annalen, 1866, 128, 100, Phil. Mag, 1866, [iv], 32, 177;
 H. Becquerel, Compt. rend., 1886, 102, 106, 103, 198, 1887, 104, 165, 777, 1691, Ann. Chim. Phys., 1888, [vi.], 14, 170, 257.
 G. H. Bailey, Brit. Assoc. Rep., 1887, p. 654, 1890, p. 778;
 Dufet, Bull. Soc. franç. Min., 1901, 24, 373

The absorption spectra of crystals of rare earth compounds are extremely rich in intense, narrow bands. As the temperature is lowered below the ordinary, the lands become even more narrow and intense 1. In the immediate vicinity of the absorption bands, the crystals exhibit the phenomenon of anomalous magnetic rotatory dispersion, aqueous solutions of rare earth salts exhibit the same phenomenon?

The absorption spectrum of a solid rare earth compound differs considerably from the spectrum of its aqueous solution. The absorption spectra of the aqueous solutions of salts of a rare earth element differ from one another, but there is a greater "family likeness" between these spectra, than between the spectra of the solid salts themselves. In fact, the dilute aqueous solutions exhibit absorption spectra which are often practically identical One or two bands observed with one salt may not be seen with another salt, and variations (usually not very pronounced) in the relative intensities of corresponding bands may occur. Moreover, the wave lengths of the "heads" of corresponding bands usually differ only by very small amounts to Thodifferences in the spectra become more pronounced, however, in concentrated solutions. In terms of the ionic theory, the results have been interpreted as indicating that absorption spectra of dilute solutions are spectra of ions, but that the absorption spectra of concentrated solutions are the spectra of bigh ions and molecules 6. This simple explanation, however, is madequate. For instance, several cases are known in which the spectra are strikingly different, instead of being practically identical as the ionic explanation requires them to be,6 and not infrequently the "heads" of corresponding absorption bands differ appreciably in position. Moreover, the ionic interpretation cannot be reconciled with the repeated observation that Beer's Law holds for moderately dilute solutions, since it leads to the conclusion that the ions of the rare earth elements produce the same absorption spectra as the molecules of their salts.5

Solutions of one and the same rare earth compound in different solvents exhibit different absorption spectra. Sometimes the differences are very striking, in other cases, the differences may be slight. For example, the absorption spectrum of neodymium chloride in glycerol bears a considerable resemblance to the spectrum of the same substance in water, the "glycerol

See J. Reequerel, Phil. Mag., 1998, (v) J. 16, 153
 J. Beequerel, loc cit.; R. W. Wood, Phil. Max., 1998, (v) J. 15, 270.; Elius, Physikal.

Zeitsch., 1909, 7, 931

J. Balla and Bursen, Annalea, 1866, 137, 1, Bursen, Popp. Annalea, 1866, 128, 100;
Phil. Mag., 1866, [iv.], 32, 177. See also G. H. Balley, tor. cit., and Rev., 1887, 20, 2769, 3325, 1888, 21, 1520, H. Becquerel, low-cit., Bettenderl, Annalea, 1899, 256, 159, 1891, 66, 143.

<sup>3325, 1888, 21, 1520.</sup> H. Becquerel, lowert., Bettendorf, Annaten, 1890, 256, 1591, 163.
43 Bahr and Bunsen, lowert., Bunsen, lowert., Buley, lowert., Becquerel, lowert.
4 See Laveing, Trans. Camb. Phyl. Soc., 1900, 18, 298. Purvis, Proc. Camb. Phyl. Soc., 1903, 12, 202, 209. Aufierld, Inaugural Insochitom (Beilm, 1904). Lamplet, Zeitzh. Phys. Rob. Chem. 1906, 56, 624. Ball. Proc. B. y. Soc., 1912, A. 87, 121.
5 Muthmann and Statzel, Rev., 1899, 32, 2053. Muthmann and Herandorf, Annaten, 1907, 355, 165. Waeguer, Ber., 1903, 36, 3055. Remerkable differences are observed, for network between the spectra of the approximation solutions of the oblighted of Mand Pr and the spectra of the approximation solutions of the chief spectra of the approximation of the corresponding double pot issum carbonates.
7 Eg. aqueous solutions of needy mum chloude and actate. The bands in the spectrum of the latter substance are less unerses, which, and tomost the red end of the servirum than of the latter substance are less unerses.

Eq. aqueous solutions of neody muon channel and act act are. The bands in the spectrum of the latter substance are less intense, wider, and no are the red end of the spectrum than those of the former (Jones and Strong, locate).
 See Rudorf, Ahren's Sammlung, 1904, No. 9, Jahresber. Elektronck, 1907, 3, 423; Zeitsch. wiss. Photochem., 1907, 5, 24, Sheppard, Photochemistry (Longmans & Co., 1914), p. 153.

bands" being, however, rather closer to the red end of the spectrum than the corresponding "water bands". On the other hand, the spectrum of neodymium chloride in ethyl alcohol is strikingly different from the preceding spectra.1

The absorption spectrum of a rare earth salt dissolved in a mixture of two solvents shows the absorption bands of the two spectra that may be observed by using the solvents separately.2 The relative intensities of these two component spectra vary with the composition of the immed solvent; in some cases several bands are observed which cannot be attributed to either of the component spectra. These facts argue strongly in favour of the view that chemical combination occurs between the dissolved salt and the solvents, as has been pointed out by H. C. Jones 3

In connection with the influence of the medium upon the absorption spectia, it is important to note that the spectra of raie earth salt solutions are considerably modified by the addition of a mineral acid, the bands (or some of the bands) usually being rendered weaker and more diffuse.4 The effect of nitric acid upon the absorption spectrum of aqueous neodymium nitrate, for instance, is very marked, the bands becoming weaker, more diffuse, and wider towards the red end of the spectrum; in this case the colour of the solution is changed in a very striking manner. The intensity of absorption of a rare earth salt is also changed, usually diminished, by the addition of another rare earth salt, whether the added salt produces absorption or not; and the various bands in the spectrum are in general unequally affected.

Absorption spectra are affected by change of temperature The nature of the change produced has been studied by Hartley, Livening, H. C. Jones and others. Rise of temperature (above the ordinary atmospheric temperature) has the effect of slightly intensifying the absorption, broadening some bands, and increasing the extent of the general absorption in the ultra-violet and infra red, the positions of the "heads" of the bands, however, are little, if at all, affected At temperatures much below the ordinary, the absorption bands become extremely narrow.7 The preceding statements may require modification at times when foreign substances are present. Thus, in the presence of calcium chloride, some of the bands in the spectrum of aqueous needymium chloride diminish in intensity with use of temperature, and the "heads" of the bands undergo a slight displacement (Jones and Strong).

It not infrequently happens that different rare earth salts present in the same solution give rise to absorption bands in almost the same region of the

H. C. Jones and others, loc cit. See also Ball, Proc. Roy. Soc., 1912, A, 87, 121.
 H. C. Jones and others, loc. cit.; cf. Miss. H. Schaefler, Physikal Zeitsh., 1906, 7.

H. C. Jones and others, loc. cit.; cf. Birss ii. Schweiner, Ingrava.
 H. C. Jones and others, loc cit; il. C. Jones, Amer. Chem. J., 1910, 41, 19; l'hil Mag., 1912, [vi], 23, 730. Zeitsch. Elektro.hem., 1914, 20, 552.
 Bunsen, Pogg. Annalen, 1868, 128, 100; l'hil. Mag., 1866, [v.], 32, 177. Lecoq de Borsbaudran and Smith, Compt. rend., 1879, 88, 1167. Soret, ibid., 1880, 91, 378, C. H. Bailey, Brit. Assoc. Rep., 1890, p. 773. Liveing, loc. cit., Strong, Physikul. Zeitsch., 1910, 11, 668; H. C. Jones and Skiong, Amer. Chem. J., 1911, 45, 1.
 Brauner, Trans. Chem. Soc., 1883, 43, 278; Demarcy, Compt. rend., 1898, 126 1039; Auer von Welsbach, Sitzunasber. K. Akad. Wiss. Wien, 1885, 92, II 317, 1903, 112, II. A, 1037; Ansaiger K. Akad. Wiss. Wien, 1905, No. 10, p. 122.
 Hartley, Sci. Trans. Roy. Dubl. Soc., 1900, II. 7, 253, Liveing, loc. cit., H. C. Jones and others, loc. cit.
 J. Becquerel, Phil. Mag., 1906, [vi.], 16, 163.

spectrum, the most striking case being that of the two bands situated at \$1690, one due to prascodymium and the other to incodymium interesting case is that of the two needymnim bands \$1610 and \$1750 and the two samarium bands A4630 and A4760. When such bands partly or wholly overlap, they may appear abnormally intense. The "heads" of such bands may also at times appear to vary in position as a composite material is fractionated.

Owing to the manner in which the absorption spectra of the rare earth compounds vary with the conditions of experiment, erroneous conclusions have not intrequently been drawn from researches upon them. Thus, Kruss and Nilson were led to suppose that the rare carth elements the compounds of which give rise to absorption spectra are in reality extremely complex substances, composed of numerous meta-elements each characterised by a single band in the absorption spectra of its com-This theory of "one band-one element" was supported at the time by Crookes, but adversely criticised by Bulley and by Schottlander 2 At the present time it cannot be seriously entertained, it will be sufficient to point out that there is no logical reason for restricting the absorption bands to the visible region of the spectrum, or for restricting the theory to the rare earth elements

Methods of Observing Absorption Spectra -- In order to observe an absorption spectrum, a suitable source of light is focussed upon the sht of the spectroscope and the absorbing medium interposed between the light

source and the sht, close up to the latter For observations in the visual region the solution may be contained in a glass test tube or, better, in a glass cell with a pan of plane, parallel ends, diffused daylight serves as a convenient source of light. In accurate work it is usually necessary to photograph the



Fig. 23 - Absorption cell.

absorption spectrum. It is then necessary, if observations are to be made for a considerable distance into the ultra violet region, to replace all glass parts in the line between sht and light source by quartz. A convenient absorption cell is shown in fig. 23, the tube A and B are joined by a broad rubber band C, and the absorbing solution poured in at D, the plates E and F which close the ends of the tubes may be made of quartz if necessary. The changes in the absorption spectrum as the thickness of the absorbing medium is varied can be easily followed by sliding the inner up and down

For photographic purposes the positive crater of a carbon arc serves as a convenient source of light so far as the visible spectrum (and particularly the red end) is concerned, for the violet and ultra violet regions, however, it is of little use, and should be replaced by a Nernst lamp Either of these sources of light gives a continuous spectrum. In order to obtain wave-length measurements, the simplest plan is generally to photograph the non are spectrum on the same plate, slightly overlapping the absorption spectrum. Some experimenters prefer to utilise a source of light that gives not a con-

<sup>&</sup>lt;sup>1</sup> Kruss and Nilson, Ber., 1887, 20, 1676, 2131, 3067, 1888, 21, 585, 2019, Kiesewetter

and Kruss, Ber., 1882, 21, 2310.

<sup>2</sup> Crookes, Chem. News, 1886, 54, 27, Bailey, Brit. Assoc Rep., 1887, p. 654; 1890, p. 773, Ber., 1887, 20, 2769, 3325, 1888, 21, 1520, Schottlander, Ber., 1892, 25, 378,

tinuous, but a discontinuous spectrum consisting of a very large number of lines, stuated close together and not differing greatly in relative intensities.

A suitable light source of this nature is obtained by sparking between carbon electrodes impregnated with the oxides of molybdenum and uranium.1

Absorption Spectra of the Rare Earths.—The absorption spectra are briefly described in the following paragraphs.2

Prascodymium. —Five absorption bands characterise the visible absorption spectrum of aqueous praseodymium chloride or nitiate, the positions of the "heads" being as follows.—

Of these bands, the violet band is broad and very intense, the blue bands are

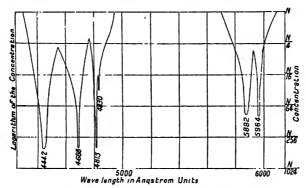


Fig. 24.—Absorption spectrum of aqueous prassodymium chloride (Rech; thickness of absorbing layer = 20 cms.).

intense, and the yellow bands weak 3 The absorption curve for aqueous prascodymum chloride, according to Rech, is shown in fig 21

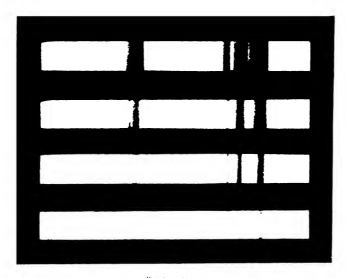
Neodymum.—The absorption spectra of aqueous neodymum chloride and nitrate are very beautiful and extremely rich in bands. The maxima that have been observed are as follows -

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7324; 6890; 6790, 6730; 6360; 6280, 6250, 6220; 5870, 5830;
5800; 5780; 5750; 5740; 5710; 5470, 5310, 5250, 5210; 5200; 5120; 5000; 4870; 4800, 4750; 1690; 4610, 4327; 4290; 4274; 4180, 3801; 3550; 3540; 3507; 3465; 3399, 3288; 3231; 3144; 3067; 3047; 3030; 3015; 2998; 2983; 2937; 2913; 2902.
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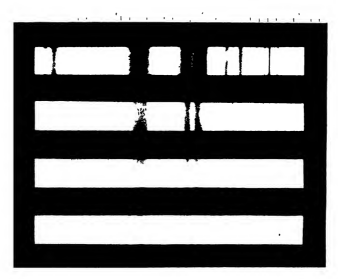
<sup>1</sup> For further information, see, e.g., Kayser, opuscit (p. 282), Jones and others, loc. cit., p. 283), Rech, Zeitsch. wiss. Photochem., 1905, 3, 411.

2 It should be mentioned that measurements of the position of a maximum of absorption by different experimenters usually differ by several Angstrom units, partly owing to experimental error and partly owing to the fact that they have not all worked with the same salt.

3 Forsling, Bihang K. Svenska Vet. Akad. Handl., 1893, 18, 1., No. 10; 1898, 23, I., No. 5; Enaglet, ibid., 1901, 26, II., No. 2, Rech, Zeitsch. wiss. Photokem., 1905, 341; Austrecht, loc. cit.; Schottlander, loc. cit.; H. C. Jones and others, loc. cit., Baxter and Stewart, J. Amer. Chem. Soc., 1915, 37, 516.



A sprain production of the consequence of the conse



 $$\rm Nr$  region  $_{\rm S}$  sorption (pectrum of a normal solution of meodynoran intrate), thickness of about nor layer (10.0, 3.3, 121, and 0.4 cms to pectively

The two bands  $\lambda 5800$  and  $\lambda 5170$  are seen when the intrate is examined, but do not appear in the chloride spectrum <sup>1</sup> The absorption curve of aqueous neodymium chloride is shown in fig. 25, which has been drawn from the data given by Rech.

Samarum.2-According to Demarcay, the following are the positions of the centres (not the "heads") of the bands in the absorption spectrum of a twenty per cent. solution of samarium nitrate in strong nitric acid (thickness of solution = 13 mm.) -

5590, 5290, 4980; 4760; 4630; 4530, 4130; 4170; 1070, 4020; 3900, 3750, 3620.

Several of these bands are complex. The positions of the "heads" of the bands in the spectrum of aqueous samarium chloride are as follows, according to Forsling

5600, 5001; 4801-4783; 4761-1727, 1632. 1413-1383, 4177; 4157; 4083, 4077, 4035-1030, 4016-1007, 3912-3932, 3906, 3752-3742; 3738-3732, 3630-3615

It is an unfortunate circumstance that the strongest of the samarium bands are very closely in the neighbourhood of prominent bands of neodymium and europium, since these are the two elements most difficult to remove from samanum.

Europeum - A neutral solution of the chloride exhibits the following maxima of absorption, the visible bands being narrow and rather faint :-

5913, 5882; 5790, 5337, **5251**; **5234**; **4656**; **4645**; 4000, 3950; 3853, 3836, 3800, 3760, 3746, 3663, **3615**; 3207, 3192, **3178**; **3168**; 2983

In the initiate solution a single band λ5913 replaces the two bands  $\lambda 5913$  and  $\lambda 5882$  , the band  $\lambda 5790$  is much more intense and narrow than in the chloride spectrum, the unsymmetrical chloride band  $\lambda 5337$  gives place to a symmetrical band  $\lambda5354$  , a single band  $\lambda5250$  replaces the two chloride bands  $\lambda5251$  and  $\lambda5234$  , and a single band  $\lambda4660$  replaces the two chloride bands  $\lambda4656$  and  $\lambda1645.^3$ 

Gadolinium.—The only absorption bands of gadolinium are in the ultraviolet region '-- 4

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3794, 3697, 3596, 3515, 3108; 3059; 3056; 3052.
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Terbium —The absorption spectrum of terbium lies in the ultra-violet with the exception of a feeble band at \$1880 :-- 5

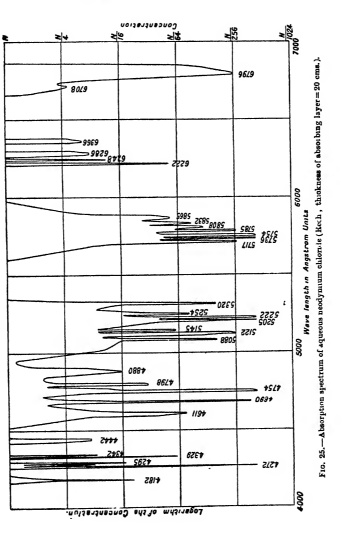
4880, 3787, 3694, 3592, 3519, 3420, 3390, 3266; 3176; 3029.

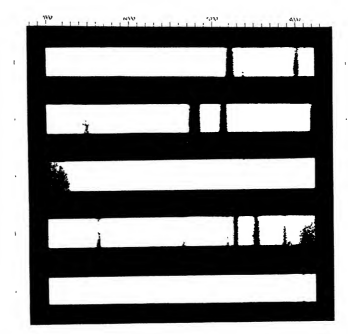
Forshing, Bihang K Svenska Vet Akad Handl., 1892, 18, L. No 4, 1893, 18, L. No
 Demarcay, Compt. rend., 1898, 126, 1039. Red., Zatsch. usw Photochem., 1905. 3,
 Holmberg, Zatsch. anorg. Chem., 1907. 53, 116; Strong, Physikal. Zettech., 1910,
 1688. Baxter and Chapin, J. Amer. Chem. Sus., 1911., 33, 1., Baxter and Woodward, 1911., 33, 270. H. C. Jones and others, los. cit., Stahl. Le Ratium, 1909. 6, 215;
 Gaimer, Arch. Set. phys. nat., 1915, [iv.], 40, 93, 199
 Forsling, los. cit.; Demarcay, Compt. rend., 1900, 130, 1185; Jones and Strong, los. cit.

Ulbain, J. Chim phys., 1906, 4, 242, cf. Demai(ay, Compt. rend., 1900, 130, 1469.

4 Ulbain, Compt. rend., 1905, 140, 1233; J. Chim phys., 1906, 4, 328, cf. Jones and Strong, loc. ctt.

<sup>&</sup>lt;sup>5</sup> Urbain, Compt. rend., 1905, 141, 521; J. Chim. phys., 1906, 4, 346.





Absorption spectra of the intrates of viriou a rice earth element, in dilute region, solution

- 1 Smarium 1 Dy prosinta nol 2 Filania Rolangia 3 Hulluna 1 mopunia

Dysprosium.—Dysprosium was originally characterised by its visible absorption spectrum, its ultra violet absorption spectrum may also be utilised for the purpose of identifying the element. The chloride solution exhibits the following maxima - 1

7530; 4750; 4515; 4275; 3970, 3865; 3795; 3650; 3510; 3380. 3225.

Holmium -This element was originally characterised by two bands,  $\lambda6400$  and  $\lambda5360$ . The positions of the absorption bands in the spectrum of aqueous holinium chloride (10 cm. thickness of N/2 solution) are, according to Holmberg, as follows -2

6577-6551; 6538-6518; **6446-6418**; **6414-6384**; 5491; 5438-5427, 5411-5401, 5385-5349; 1910, 1860-1846, 1830, 1683-1672; **4554-4459**; 4223-4216; 4179-4152

Erbium. -Owing to the difficulty of preparing erbium salts free from impurities which give absorption spectra, the spectrum of erbium is not known as exactly as could be desired. From the work of Forsling 3 the erbium spectrum appears to exhibit maxima at the following positions -

6665, 6530-6515; 6483, 5490, 5412, 5231; 5202-5188, 4908; 4871; 4815, 4534, 4497; 4427; 1081, 1069, 4058, 1050, 3792, 3641, 3637, 3588; 3561.

This list comprises all the maxima given by Urbain,4 but contains three maxima (\lambda 6483, \lambda 5490, \lambda 4531) that could not be observed by Holmann and Burger.5

Thultum.—The following are the positions of the maxima in the absorption spectrum of thuhum -6

7015; 6845-6828; 6590: 4640; 3600.

Reflection Spectra.—It has already been mentioned that the absorption spectra of solid rare earth compounds may be observed by allowing ordinary white light to fall upon them and examining the reflected light spectroscopically. This method is very useful in cases where, owing to the fact that the substances are opaque, the absorption spectra cannot be observed by transmission. Absorption spectra observed by reflection are usually referred to as reflection spectra

The question naturally arises as to whether the absorption spectrum of a substance observed by transmission is, or is not, identical with the absorption spectrum observed by reflection. So far as the rare earths are concerned,

<sup>&</sup>lt;sup>1</sup> Urbain, Compt. rend , 1906, 142, 785 , Lecoq de Boisbaudian, ibul , 1886, 102, 1008,

Urbain, Compt. rend., 1906, 142, 785, Lecoq de Boisbaudian, that, 1886, 102, 1003, 1005; see also Jones and Strong, tor cit.
 Hölmberg, Arkiw Ken. Min Geol., 1911, 4, Nos. 2 and 10, Zeitsch. anong. Chem., 1911, 71, 226, cf. Urbain, J. Chim. phys., 1906, 4, 31; Soret, Compt. rend., 1879, 89, 52; Clove, that, 1879, 89, 478, 708, Langlet, Arkiw Kem. Min. Gool., 1907, 2, No. 32; Forsling, Bihang K. Stenska Vet.-Akad Handl., 1902, 28, II., No. 7.
 Urbain, J. Chim., phys., 1906, 4, 31.
 Hofmann and Burger, Ber., 1908, 44, 308; see also Hofmann, Ber., 1910, 43, 2631; Jones and Strong, loc. cit., Puivis, Iroc. Camb. Phil., Noc., 1903, 12, 202, 206; Langlet, Zeitsch. physikal Chem., 1906, 56, 624.
 Forsling, Bihang K. Stenska Vet.-Akad Handl., 1899, 24, I., No. 7; see also Cleve, Compt. rend., 1879, 89, 419, 478, 1880, 91, 381, Thalen, thid., 1880, 91, 45, 326, 376; Urbain, J. Chim. phys., 1906, 4, 81.

there is no experimental work available from which an answer may be deduced, but from work on other substances it has been found that the spectra are different. Theoretically there should be a difference, because the reflecting power of an absorbing medium increases with the refractive index, and owing to the phenomenon of "anomalous dispersion" being exhibited in the neighbourhood of an absorption band, it follows that an absorption band in the reflection spectrum should be relatively less intense on the red side and more intense on the blue side than the corresponding band in the absorption spectrum by transmission 1

The reflection spectra of the rare earths were discovered by Bahr and Bunsen, and have been subsequently studied by Hartley, Crookes, Humpidge, H. Becquerel, Dennis and Chamot, Exner, Haitinger, and others <sup>2</sup> Anderson <sup>3</sup> and Joye 4 have described convenient methods for photographing the spectra.

Examination of the reflection spectra of the oxychlorides of the rare earth metals is of considerable value for following the course of separation of these metals by fractional crystallisation, etc. Only a few milligrams of material are necessary 5 Instead of the oxychlorides the basic intrates may

For the rapid examination of the reflection spectrum of a small quantity of a solid, the light of the carbon are is concentrated upon it. Between the illuminated solid and the sht of the spectroscope a lens is placed to focus the light from the solid on to the slit. It is possible in this manner to observe absorption bands in the spectrum of a substance when its aqueous solution no longer appears to exhibit selective absorption 6

For detailed descriptions of reflection spectra the reader is referred to the original memons.7

## FLAME SPECTRA.

When the oxides, borates, or phosphates of some of the rare earth elements are heated to meandescence in a non-luminous flame, such as that of a Bursen burner, the light contted is found to give not a continuous spectrum, as would be anticipated, but a sharply discontinuous band spectrum. The

<sup>&</sup>lt;sup>1</sup> See R. W. Wood, Phil May, 1902, [v1], 3, 620.

<sup>2</sup> Bahn and Bunsen, Annalon, 1866, 137, 1, Hattley, Trans. Chem. Soc., 1882, 41, 210, Chem. News, 1886, 53, 179, Crookes, Compt. rend., 1886, 102, 506, Humpedge, Chem. News, 1886, 53, 154, H. Beequerel, Ann. Chem. Phys., 1888, [v1], 14, 170, 257; Hattinger, Monatsh., 1891, 12, 362, Dennis and Chamol, J. Amer. Chem. Soc., 1897, 19, 799; Exner, Silkanayber, K. Akad. Wiss. Wien, 1899, 108, Ha, 1252, Urbain, vide. infra., Waegner, Ber., 1903, 36, 3055; Zeitsch unorg. Chem., 1904, 42, 122, Muthman and Hersundel, Annalon, 1907, 355, 165, Anderson, vide. infra., Holmann and Hoschele, Ber., 1908, 41, 3783, Holmann and Kunmenther, inde. infra. Holmann and Hoschele, Ber., 1914, 47, 240, Schaum and Wustenfeld, Zeitsch. wiss. Photochem., 1911, 10, 226, Joye, vide. virta.

Anderson, Astrophys J., 1907, 26, 73
 Joye, Arch See phys nat, 1913, [w], 36, 41
 Urbain, Ann Chim Phys, 1900, [vn], 19, 222. For the rapid preparation of the

<sup>\*\*</sup> Othann, Ann. Chim. Phys., 1900, [vn.], 19, 222. For the rapid preparation of the oxychlorides, see p. 255

\*\*O'Urbain, Inc. id.\*\*

\*\*T See Joye, Arch. Sci. phys. nat., 1913. [iv.], 36, 41 (Nd<sub>2</sub>O<sub>1</sub>, Nd<sub>2</sub>O<sub>2</sub>, 3H<sub>2</sub>O, 2Nd<sub>2</sub>O<sub>3</sub>, 3H<sub>2</sub>O, Nd<sub>2</sub>O<sub>4</sub>, 11,O, Nd<sub>2</sub>O<sub>4</sub>O<sub>4</sub>), Nd<sub>3</sub>O<sub>4</sub>, NdCl<sub>3</sub>, NdCl<sub>3</sub>, NdBl<sub>3</sub>, Nd<sub>2</sub>C(O<sub>4</sub>), Nd<sub>3</sub>C(O<sub>4</sub>), Nd<sub>3</sub>O<sub>4</sub>, Nd(NO<sub>4</sub>), Oll<sub>2</sub>O<sub>5</sub>O<sub>4</sub>, 1913, [iv.], 36, 431. (Pr<sub>2</sub>O<sub>4</sub>, Pr<sub>2</sub>O<sub>3</sub> 3H<sub>2</sub>O, Pr<sub>4</sub>O<sub>4</sub>, Pr<sub>4</sub>O<sub>4</sub>, Pr<sub>4</sub>O<sub>5</sub>O<sub>4</sub>), 10H<sub>2</sub>O<sub>5</sub>O<sub>4</sub>, 10H<sub>2</sub>O<sub>5</sub>O<sub>4</sub>, 10H<sub>2</sub>O<sub>5</sub>O<sub>4</sub>, 10H<sub>2</sub>O<sub>5</sub>O<sub>4</sub>, Er<sub>2</sub>O<sub>5</sub>O<sub>4</sub>, Er<sub>2</sub>O<sub>5</sub>, Er<sub>4</sub>(SO<sub>4</sub>)<sub>3</sub>, Er<sub>2</sub>O<sub>5</sub>, ErCl<sub>3</sub>); Hofmann and Hoschelo, Ber., 1914, 47, 240 (PrOCl, NdOCl, SmOCl, ErOCl).

spectrum of neodymia was discovered by Bunser 1 in 1864, and that of cibia by Bahr 2 m 1865. Up to the present time it has been found that, under suitable conditions, prascodymia, neodymia, samaria, erbia, and thulia? give use to these emission spectra. No satisfactory explanation of their production has yet been advanced. It has been stated that the spectra are due to thin layers of glowing vapour," but this is almost certainly not the case 9

It is not definitely known whether a pure rare earth gives rise to a flame spectrum or not. It is stated that prascodymia gives no such spectrum unless mixed with another oxide, eg alumina or magnesia, while the flamo spectrum of neodymia alone is said to be very hazy. Most observations of these spectra have been made with erbia, and pine erbia has not yet been prepared.

The flame spectrum of a rare earth differs considerably from its cathodic phosphorescence spectrum, but is closely connected with its reflection spectrum. The emission bands in the flame spectrum of crbia, for instance, are in almost exactly the same positions as the absorption bands in the reflection spectrum of the same substance. The reflection spectrum varies with the temperature, and in the cases of erbia and neodymia it has been found by Anderson that the bands in the flame and reflection spectra occupy identical positions when examined at the same temperature. This result is in accordance with Knelhoft's Law  $^{10}$ 

The flame spectra of different compounds of the same rare earth element exhibit a marked resemblance to one another, without, however, being identical 11 Morcover, it follows from what has been already stated that these emission spectra must correspond approximately with the absorption spectra of the compounds of the same rare earth element. This correspondence was noted by Bahr and Bursen

The flame spectry of the rare earths are of little or no practical value to the chemist

# REVERSION SPECIFIC

When the spark spectrum of a solution of a salt is examined in the usual way (p. 305), the external inctallic electrode and the salt solution are

- Bunsen, Annalea, 1864, 131, 255. Bahr and Bunsen, ibid., 1866, 137, 1.
   Bahr, Innalea 1865, 135, 376. Bahr and Bunsen, ibid.
   Hartinger, Monotch., 1881, 12, 362.
   Amer von Welsbach. Set warder K. Akad. West. Wen., 1885, 92, 11–317. Memotch., 1885, 6, 477. Hartinger, ibid. etc., Schaum and Wustenfeld, Zeitsch. west. Photochem., 1911, 10, 226. Anderson, Isti. phys. J., 1967, 26, 73.
   Levoq de Borsbaudran, Compt. vend., 1879, 88, 322. 89, 212. Bettendorf, Annalen, 1891, 263, 164.
   Bahr, ibid. etc., d., Bahr and Bunsen, ibid. d., Levoq de Borsbandran, Compt. vend., 1873, 76, 1939. Spectres tummente (Paris, 1874). Thaken, Compt. vend., 1886, 92, 226. Crookes, ibid., 1888, 102, 506. Hofmann and Bugge, Ibid., 1998, 41, 3783. Hofmann and Kunmenther, Zeitsch. physikal. Chem., 1910, 71, 312. Anderson, Ibid. etc., Schaum. and. Wustenfeld, Ibid. etc.

- Zettect physikal, Chem., 1910, 71, 312. Anderson, the cit., Schaim and Wastenfeld, loc. cit.

  7 Thalen, Campt. rend., 1880, 91, 376.

  4 Huggins, Proc. Roy. Soc., 1870, 18, 546., Thil. May., 1470. [iv.], 40, 302., Reynolds, thid., 1870, [iv.], 40, 302. Reynolds, 4 Anderson, Istraphys. J., 1907, 26, 73.

  10 Anderson, loc. cit., see also Schaim and Wustenfeld, loc. cit., and Heimann and Kunneuther, loc. cit.

  11 Leave de Bushandry, Compt. 2010. 1873, 76, 1080.
  - 11 Lecon de Boisbaudran, Compt rend., 1873, 76, 1080.

connected to the positive and negative poles respectively of the induction coil. If this arrangement is reversed, so that the solution is positive to the external electrode, the line spectrum of the metal can scarcely be seen, but with solutions of samarum, curopium, terbium, and dysprosium salts, if a long spark is used, the liquid surface becomes phosphorescent immediately below the external electrode (see fig 26). The phosphorescent light, when examined spectroscopically, is found in each case to yield a discontinuous

band spectrum. These spectra are known as reversion spectra (spectres de renversement) They were discovered by Lecoq de Boisbaudran 1

The reversion spectra bear a striking resemblance to the corresponding eathodic phosphorescence spectra In dilute solution they are practically independent of the acid radicle present. The intensity of a reversion spectrum is approximately proportional to the concentration of the solution, and its sensitiveness is comparable with that of an absorption spectrum 2

Samarrum chloride or nitrate solution gives a yellow phosphorescence the spectrum of which consists of three bands, the maxima being at  $\lambda 6440$ ,  $\lambda 6000$ ,  $\lambda 5640^3$ 

Europeum (or Zz) gives rise to a red phosphorescence Fig. 26 -Apparatus for A dilute acid solution shows two beautiful bands, production of rever- (1)  $\lambda6160$  to  $\lambda6110$ , maximum at  $\lambda6125$ , and (ii )  $\lambda5970$ the phosphorescent to \$5920, maximum at \$5890 4

Terbum (or Z<sub>g</sub>) gives use to a beautiful given phosphorescence. The spectrum of the chloride solution shows four distinct bands, the maxima being situated at λλ6205, 5858, 5432, and 1870 5

Dysprosium (or  $Z_a$ ) gives rise to a yellow phosphorescence the spectrum of which consists of two bands. The maxima are at \$5730 and \$1765.0

### CATHODIC PHOSPHORESCENCE SPECIFIA.

Historical.-In 1881 Crookes discovered that numerous substances, when exposed to the bombardment of the cathode rays, emit a phosphor-

Lecoq de Borsbaudran, Compt. rend., 1885, 100, 1437; Chem. News, 1885, 52, 4
 The presence of certain elements, e.g. non- and inthenium, inhibits the production of the reversion spectra (Lecoq de Borsbaudran, Compt. rend., 1886, 103, 113)
 Demarqay, Compt. rend., 1900, 130, 1185, cf. Lecoq de Borsbaudran, chal., 1892, 114, 575; 1893, 116, 611
 Uthain, J. Chim. phys., 1906, 4, 247, Demarqay, Compt. rend., 1900, 130, 1169, Lecoq de Borsbaudran, Compt. rend., 1892, 114, 575, 1893, 116, 611, 674.
 Lecoq de Borsbaudran, Compt. rend., 1886, 102, 899, Urbain, J. Chim. phys., 1906, 4, 348

<sup>Lecoq de Borsbaudian, Compt. rend., 1886, 102, 899, Ulbam, J. Chim. pags., 1200, 4, 348
Lecoq de Borsbaudian, Compt. rend., 1886, 102, 899, Ulbam, thal., 1906, 142, 785
For further information concenning reversion spectra, see the relatences to Lacoq de Borsbandian cited on p. 290, and Urbain, Ann. Chim. Phys., 1909, [vin.], 18, 222
The work of Cipokes on phosphorescent spectra will be found in the following memons or reprints: Compt. rend., 1379, 88, 283; 1881, 92, 1281, 1885, 100, 1380, 1495, 1886, 102, 506, 646, 1464, Phil. Trans., 1883, 174, pt. 111., 891, 1885, 176, pt. 11., 691; Proc. Roy. Soc.. 1881, 32, 256, 1883, 33, 262; 1885, 34, 414, 1886, 40, 77, 236, 602, 1887, 42, 111; Chem. Neus, 1881, 43, 237; 1883, 47, 261, 1884, 49, 159, 169, 181, 194, 205; 1885, 51, 301; 1886, 53, 75, 133; 54, 13, 29, 38, 40, 54, 63, 76, 155, 1889, 60, 27, 39, 51, 63; Brit. Assoc. Reports, 1886, p. 558, Trans. Chem. Soc., 1888, 53, 487, 1889, 55, 250.</sup> 

escent light which, analysed spectroscopically, shows in general a faint continuous spectrum with a more or less decided concentration in the region corresponding with the colour of the emitted light. Less commonly, the spectrum is discontinuous.

In studying these discontinuous spectra, Crookes was particularly interested in a bright citron band at  $\lambda5740$ , and endeavoured to identify the element which gave rise to it. He assumed a priori that the brightness of the band  $\lambda5740$  would increase in proportion as the element responsible for it concentrated in his preparations. In general he decomposed his materials with sulphuric acid, evaporated off the excess of acid, and exposed the anhydrous sulphates to the cathode rays. After many experiments, the element was traced to the yttrium group of the rare carticlements. Accordingly, he prepared specimens of the yttria cartis in as price a state as was possible at that period, examined their sulphates, and identified the sought for element with yttrium. In this, however, he was wrong, for, as will be seen later, the citron band is due to dysprosium.

A pair of orange bands, \$5970 and \$6030, also attracted attention, and an endeavour was made to trace the element X which produced them. As the result of many experiments carried out with this end in view, Crookes discovered the remarkable fact that the compounds of X are only sensibly phosphorescent when intimately mixed with certain other substances, of which the compounds of calcium are at least as good as any, and consequently he nearly always introduced calcium sulphate into his materials when studying their phosphorescence spectra. As before, he traced X to the rare earth elements, and on examining specimens of Cleve's rare earth preparations, correctly identified the element X with

Still-another band, a red one at  $\lambda 6090$ , attracted his attention. Sometimes it was nebulous, at other times it was clear and sharp, and almost as narrow as a spark line. In the phosphorescence spectrum of samarium sulphate it was very feeble, it did not occur in the spectrum of yttrium sulphate, but in the spectra of certain mixtures of these sulphates it stood out brilliantly. Grookes called it the anomalous line, and originally supposed that it corresponded to a vibration that could only be set up by the simultaneous presence of samarium and yttrium. Shortly afterwards, however, he abandoned this idea, and attributed the anomalous line to a separate element which provisionally received the designation  $S_{\delta}$ . It will be seen subsequently that  $S_{\delta}$  is the element enoquium.

('rook's observed that the entron band was accompanied by numerous red, green, blue, and violet bands, a number of which were extremely sharp and narrow. In 1886 he discovered that these bands varied among themselves in relative intensity when vitria was methodically fractionated, and concluded that yttimin must be an extremely complicated substance which can be split up, at least partially, into numerous simpler substances, each characterised by a phosphorescence spectrum of great simplerty, consisting for the most part of a single band. From further experiments, he drew the conclusion that samarium and Marignac's Y<sub>o</sub> (gadoliminin) are also complex substances. The simpler constituents of these "elements" he called meta-elements and gave provisional names, and he put forward a new theory of the nature of the chemical elements in his British Association address in 1886. Owing to the striking and original nature of these views, they created great interest.

The compositions of samarum, yttrum, and gadolmium in terms of their component meta-elements were given by Crookes in 1889 as follows -

| Meta-<br>element                             | λ                            |                     | Mcts-   A                                                                                                    | Meta-clement   A                                                                                                                       |
|----------------------------------------------|------------------------------|---------------------|--------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------|
| Så<br>G,<br>G <sub>y</sub><br>G <sub>g</sub> | 6040<br>5970<br>5640<br>6390 | <br>  Sansarium<br> | $\begin{pmatrix} G_{\beta} & i \frac{5450}{6190} \\ G_{\zeta} & i \frac{6190}{6190} \end{pmatrix}$ Gadolmium | C <sub>α</sub> 4820<br>C <sub>β</sub> 5170<br>C <sub>ζ</sub> 6190<br>C <sub>δ</sub> 5740<br>C <sub>γ</sub> 5970<br>C <sub>γ</sub> 6470 |

It will be seen that a meta element could enter into the composition of more than one "element"

These ideas were all opposed by Lecoq de Borsbandran, who in 1885 discovered two new rare earth elements,  $Z_a$  and  $Z_b$ , belonging to the terbia group, by means of their reversion spectra, and was struck by the resemblance between these spectra and the phosphorescence spectra attributed by Crookes to  $G_{\delta}$  and  $G_{\beta}$  respectively. By careful fractionation, the French spectroscopist prepared pure yttria, and found it to be devoid of phosphorescence, the addition of a little  $Z_{\alpha}$  and  $Z_{\beta}$ , however, produced the "yttrium" spectrum described by Crookes He further found that whereas the atomic weight of yttrum is about 89, the values for  $Z_a$  and  $Z_B$  are about 160, and that the spark spectra of these elements show no lines that can be attributed to yttrium - Lecoq de Boisbaudran therefore concluded that pure substances are not phosphorescent, and that yttma acts merely as a diluent, in which  $Z_a$  and  $Z_B$  are able to exhibit their phosphorescence spectra, in precisely the same way as Crookes had shown lime to act as a diluent towards samarinin He further showed the correctness of this by studying numerous phosphorescent systems made up of compounds of some of the common elements. The views propounded by Lecoq de Buisbaudian were not, however, accepted by Crookes, and the Crookes—de Boisbaudian controversy served but to emphasise the difficulties of rare earth chemistry without either of the protagonists succeeding in establishing his views to the satisfaction of chemists in general In 1900 Muthmann and Baur 2 were led to beheve that Lecoq de Boisbaudian's conclusions were incorrect, in the following year, however, Baur, in association with Marc, arrived at the opposite conclusion 3

At the present time the controversy may be regarded as closed beautiful researches of Urbain and his collaborators have clearly demonstrated the correctness of Lecoq do Borsbaudran's views, and the fasemating theory of the meta-elements is now seen to be nothing more than a daring

<sup>Lecoq de Borsbaudrau's work on phosphorescence spectra will be found in the following papers: Compt. rend., 1885, 101, 552, 588; 1886, 102, 153, 1536, 103, 113, 468, 627, 629, 1064, 1107, 1887, 104, 330, 478, 554, 824, 1581, 1680; 105, 45, 206, 258, 261, 301, 301, 301, 347, 784, 1228; 1888, 106, 452, 1386, 1781, 107, 311, 168, 490, 1890, 110, 24, 67.
Muthiman and Baur, Rev., 1900, 33, 1748.
Baur and Marc, Rev., 1901, 34, 2460
Urbain, Ann. Chim. Phys., 1909, [viii.], 18, 222-376, Bruninghaus, thid., 1910, [viii.], 20, 519-546, 21, 210-283
See also Urbain, Compt., rend., 1906, 142, 205, 1518, 143, 229; 1907, 145, 1335, J. Chim., phys., 1906, 4, 233, Urbain and Scal., thid., 1907, 144, 1368</sup> 

but erroneous interpretation of experimental results. By its fall the chemistry of the rare earths lost some of its charm, but it has gained enormously in simplicity  $^{1}$ 

General Laws.-Experiment shows that pure substances are not sensibly phosphorescent when exposed to the exthode rays, the phosphorescence exhibited by a "pure" substance is found to be attributable to the presence of traces of impurities

The addition of a trace of a second compound B to a pure non-phosphorescent compound A frequently results in the production of an extremely phosphorescent mixture? The manner in which the intensity of phosphorescence varies as the relative amounts of A and B are changed has been examined by Urbain and Braninghaus for a large number of mixtures, and they have found that as the amount of one compound, say B, is varied from an exceedingly small percentage upwards, the intensity of the phosphorescence (the mode of exeitation being the same throughout) quickly increases to a maximum and then steadily diminishes <sup>4</sup>. The percentage of B present in the nuxture of maximum phosphoreseing power depends upon the natures of both A and B, but is always of the order of 0.5 to 10 per cent, and unixtures with 20 per cent, and more of B are often not sensibly phosphorescent 1

When B is a rare earth compound, the phosphorescence spectrum is sharply discontinuous, often exhibiting a large number of quite narrow bands, and as the percentage of B alters, so also does the colour of the phosphorescent light Spectroscopic abservation shows, however, that the preceding "maximum law" holds good for each band in the spectrum. The colour changes are due to the fact that for mixtures of two given substances A and B, the luminous maxima of the bands correspond to different percentages of B, in other words, the bands vary independently. Concerning the maxima of the various bands, the general rule may be stated that the most refrangible bands are at their maxima for smaller percentages of B than is the case with the least refrangible bands. Hence, as the percentage of B changes from zero upwards, the changes observed in the colour of the phosphorescent light occur in the direction from blue to red along the spectrum

The percentage of B corresponding to the luminous maximum is independent of the intensity of the eventing cathode rays, and also of the temperature, though the latter factor may produce sensible changes in the phosphorescence spectrum.

. . --- - --

<sup>&</sup>lt;sup>1</sup> In addition to the memors aloody cited, the following also relate to phosphorescence spectra: Mangna, (Farres completes 1962, vol. n. p. 811), Bettendort, Annalen, 1891, 263, 164, 1892, 270, 376. Marc, Ber. 1906, 39–1392, Wiedemann and Schmelt, Antsch., physikal Chem., 1895, 18, 529. Goldstein, Set empler K. Aland Wies, Berlin, 1900, p. 821. Cimpbell Swinton, Prec. Roy. Soc. 1899, 65, 115. Kowalski and Garmer, Complemental 1907, 144, 836.

p. 821. Complet! Swinton, Prec. Ron. Soc. 1999, 05, 115. Rowarski and Garmer, Completend, 1907, 144, 836.

It may be stated here once for all that mere mechanical mixing is not sufficient, substances must be extremely intimately mixed by chemical processes. Thus the rare earth oxide mixtures reteined to late are mode by precipitating the oxalates together from solution, washing, drying, and igniting the mixture. By treatment with concentrated sulphure acid, sulphate mixtures are obtained, etc.

This law is called la lor de Poptimium by its discoverers.

It is a forest to the state of the conden law in the greatly and that which expresses the

In this passes search to the treatment of the properties of the search and that which expresses the manner in which the light emitting power of a maximize of thoma and certa varies with the composition, the maximum here corresponds to 0.9 per cent of certa, and the light-emitting power diminishes rapidly as the percentage of certa is decreased of increased from this value

The preceding observations naturally suggest the question as to whether the phosphorescence spectrum observed in a binary mixture is to be attributed to B, which in the mixture of maximum phosphorescing power does not exceed 1 per cent of the whole, or to A, or to both 1s, for example, the phosphorescence spectrum of terbia—gadolinia mixtures to be attributed to terbia, to gadolinia, or to both? It is found that mixtures of terbia and either yttria, line, alumina, or gluena give phosphorescent spectra which, without being identical, exhibit a very strong family resemblance to one another and to the terbia—gadolinia spectrum—Morcover, the five mixtures phosphoresce most brilliantly for quantities of terbia of about 1 per cent. Gadolinia, however, when mixed with any of the other four oxides, yields a mixture which produces no visible phosphorescence—The observed phosphorescence of the system terbia—gadolinia must therefore be attributed to terbium, in just the same way as the absorption spectrum of an aqueous solution of neodynnum chloride is attributed to the neodynnum chloride.

Accordingly, in a system A—B which exhibits a maximum phosphorescence when the amount of B present is only of the order of 1 per cent, the phosphorescence is a specific property of B. The substance B is called the phosphorogen (or active matter) and A the diluent

It has been mentioned that the spectra of terbium in various diluents exhibit a marked similarity, but that they are not identical. This influence of the diluent is a general phenomenon, and was observed by Crookes, particularly for the samarium spectrum. The spectra of a phosphorogen in various diluents exhibit bands in the same regions of the spectrum, but the positions of similar bands are not identical. There is very little difference, however, in passing from one diluent to another if the metallic constituents of the diluents are the same, e.g. the spectra of samarium are very similar when the oxide, sulphate, fluoride, and borate of samarium are diluted in the corresponding salts of calcium, but a change in diluent from, say, lime to yttira, produces a small but decided change in the positions of the bands. The nature of the spectrum of a phosphorogen also depends upon the physical state of the diluent, and hence, in the oxide systems to be described later, differs somewhat according as the preparation has been made by heating the oxidates or the sulphates.

It is beyond the scope of this book to enter into a discussion of the various theories of phosphorescence, but it may be observed that phosphorogens are coloured substances, compounds of metals the salts of which are coloured and exhibit absorption spectra in solution. Good diluents, however, are colourless compounds practically devoid of electrical conductivity. These statements apply to systems which are visually phosphoroscent, it is possible for a colourless substance to act the rôle of a phosphorogen, gadolima being a case in point, but its phosphorescence spectrum lies in the ultra-violet region. It is significant that gadolimium exhibits a well-defined ultra-violet absorption spectrum. Apparently, a substance does not act both as a diluent and a phosphorogen, but does act both as an absorber and a phosphorogen in the same region of the spectrum. However, in the case of the are earths at least, the phosphorescence and absorption bands do not coincide

In practice, systems more complex than the binary systems hitherto discussed are encountered. Various ternary systems, in which two phosphorogens are diluted in a common diluent, have been studied by Urbain

<sup>1</sup> On this point see Bruninghaus, loc cit.

and Scal, and it has been found that the observed phosphorescence spectrum of such a system is made up of the spectra of the two phosphorogens, though in each component spectrum the relative intensities of the bands may differ somewhat from those observed in the spectrum of the corresponding binary system. For each spectrum the low de Poptimum holds good, but each phosphorogen appears to diminish the luminosity of the spectrum of the other, and in this action the less intense phosphorescence suffers much more than the other.

The Phosphorescence Spectra of the Rare Earths. So far as visible phosphorescences are concerned, lanthana, gadohnia, and yttira act as dilnents, and prascodyma, neodyma, samaria, europia, terbia, dysprosia,

and crbia act as phosphorogens, and rare earth mixtures in which a large excess of diluent is present phosphoresce brightly under the influence of the cathoderays. It is curious to notice that hobina and thulia do not appear to act as phosphorogens, although the safts of bolimium and thulium exhibit well defined absorption spectra.

The form of vacuum tube best smited to the observation of cathodic phosphorescence spectra is indicated in fig. 27. The electrodes A and C are made of aluminum. The substance inder examination is placed in the lower part of the apparatus (which is joined to the upper part by a lubricated ground glass joint) with its surface slightly inclined. The upper part of the apparatus is scaled on to a merciny pump by means of a side tube not shown in the diagram. For observing the spectra, a spectroscope of not too great dispersion should be used, and the slit opened fainly widely. In photographing the spectra, an exposine of trem twenty minutes to two hours is necessary, and in order to include the ultra yield region, a special form of tube with a

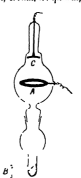


Fig. 27 —Apparatus for producing cathodic phosphorescence

quartz window B should replace the bottom part of the apparatus (see the

dotted portion of fig. 27)

BINARY SYSILMS <sup>1</sup> From the practical point of view, systems in which lanthana is the diluent are not of much interest, those with gadolina or yttria are much more important.

 $Pracodymium -- \mbox{When diluted in lime, pracedymaa exhibits a red phosphorescence, and so also does the sulphate when diluted in calcium sulphate. The spectra are indicated in figs. 28 and 29 <math display="inline">^2$ 

The relative intensities of the lands in the spectra are represented in the accompanying diawings—by plotting the intensities as ordinates against—a scale of wave lengths (unit = 10 angstroms)—They represent Uliabin's results—and, indees otherwise stated, refer to the spectra when at their luminous maxima (about 1 per cent of phosphologen)

To prepare binary rate earth mixtures, propriete the oxalates together, and calcine in a buille lumace, to prepare rate earth—line maximes, precipitate the mixed carbonates, and ignite to oxides over a good Bunson binner, to prepare rate earth—alumina or rate earth—flienna mixtures, strongly grate the precipitated moved hydroxides, etc. The dihents should be pure. Ordinary "part This gives an otang phosphorescence the to traces of manganese, calcium nitrate should be fractionally ray-tallised from intricated in order to eliminate the manganese (in the tails). Ordinary alumina gives a red phosphorese nea due to traces of chromium; the intriate should be fractionally crystallised from intricated to remove the chromium (in the tails), etc.

2. The relative intensities of the bands in the spectra are represented in the accompanying drawings by relating the intensities as ordinates against a scale of wave lengths (units).

Neodymium.—The phosphorescence is not very bright. With the system  $Nd_2O_8$ —CaO, the following maxima are observed —

5045; 4690; 1660, 4610, 4575; 4515, 4400, 4295; 1270, 4230; 4220; 4190; 3980; 3920.

The system  $\operatorname{Nd}_2(SO_4)_3$ —CaSO<sub>4</sub> shows only one broad band, from  $\lambda 5770$  to  $\lambda 5010$ .

Samarum —The phosphorescence produced by samarum is orange-yellow, orange, or orange-red in colour. The spectra observed when the diluents are

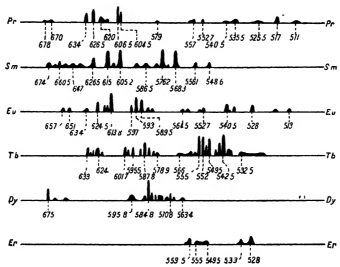


Fig. 28 -Phosphorescence spectra of the rate out his diluted with quicklime.

calcium oxide, sulphate, and fluoride and gadolimium oxide and sulphate are shown in figs 28-32.

Europium—The colour of the europium phosphorescence is generally orange-red or camine red. The spectra observed with oxide, sulphate, and fluoride of calcium and oxide and sulphate of gadolina as diluents are shown in figs. 28–32

The optimum mixture for  $\operatorname{Eu_1O_8} - \operatorname{Gd_2O_8}$  contains only 0.42 per cent, of europia. With this system, the variations in the relative intensities of the bands with the dilution is exhibited in a most striking manner.

Europium was shown by Demarçay¹ to be identical with Crookes's Ss. The identity has been disputed by Crookes,² since the position of the "anomalous band" given by Demarçay and later by Urbain does not agree

Demarçay, Compt. 2cnd., 1901, 132, 1484
 Chookes, Chem. News, 1901, 84, 2; 1905, 92, 25, Proc. Roy Soc., 1905, A, 76,

with the position \$\lambda 6090\$ that he found. This discrepancy, however, is due to change of the diluent 1

Gadolinium —The phosphorescence spectrum of gadolinium has wholly in the ultra-violet region, and for the system Gd<sub>2</sub>O<sub>3</sub>—CaO comprises the following maxima -

3166, 3159; 31555, 3153; 31505, 3147; 3144, 31405; 3138, 3136; 3134; 3130, 3091, 30885, 5085, 28283, 27895

The gadolimum spectrum is of particular interest. In 1899 Crookes<sup>2</sup> observed a new phosphorescence spectrum in the ultra-violet region and

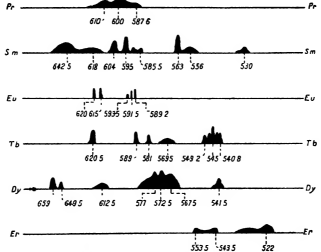


Fig. 29.—Phosphorescence spectra of rare earth sulphates diluted with calcium sulphate

attributed it to an element victorium.3 According to Crookes, "victoria" was more basic than terbia, but less so than yttria, and the atomic weight of "victorium" was equal to about 117

In 1905 Crookes concluded that Urbain's purest gadolinia contained "victorium" as an impurity - Urbain has shown, however, that the phosphorescence spectrum which defines "victorium" is really part of the gadohinum spectrum? If "victorium" were present merely as impurity in gadolinium, then dilution of gadolinia with a large excess of lime should cause the "victorium" phosphorescence spectrum to vanish. Actually, it greatly intensifies the spectrum, which is shown very feebly by practically pure gadolinia. The

<sup>1</sup> On this point see also Lecoq de Borsbaudian, Compl. icnd., 1892, 114, 575, 1893,

<sup>116, 611.

- (</sup>Crookes, Proc. Roy Soc., 1899, 65, 237, Chem. News, 1899, 79, 212; 80, 49

5 First called monitum

4 Crookes, Chem. News, 1905, 91, 61, 1906, 93, 143

6 Urbain, J. Chim phys., 1906, 4, 321, Ann Chim. Phys, 1909, [viii.], 18, 301

"victorium" spectrum, in fact, is at its maximum in the system  $\mathrm{Gd}_1\mathrm{O}_3$ — $\mathrm{CaO}$  when about 1 per cent, of gadolima is present. According to Urbain, the method employed by Crookes in isolating "victorium" can only lead to the production of a mixture composed essentially of gadolimina and yttrium, the number of fractions being far too small to lead to the isolation of a new earth. Moreover, there is no corroborative evidence in favour of the view that a new element of atomic weight 117 can be separated anywhere between samarium and dysprosium. Most convincing of all, a mixture of 226 parts of gadolima and 58 parts of yttria corresponds to an "atomic weight" of 118 and possesses all the properties associated by Crookes with oxide of "victorium". The "victorium" spectrum observed by Crookes is, then, the spectrum of gadolimium in the system  $\mathrm{Gd}_2(\mathrm{SO}_4)_3$ — $\mathrm{Y}_2(\mathrm{SO}_4)_3$ 

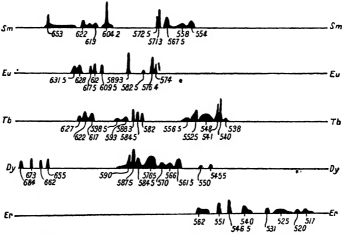


Fig. 30.--Phosphorescence spectra of the rare earth finorides diluted with calcium finoride.

The spectrum of gadolimium is more sensitive in calcium sulphate than in calcium oxide

Terbium—The phosphorescence of a system containing terbium is blue, bluish-green, or green, according to the amount of terbium present. The spectra of terbium in calcium oxide, sulphate, and fluoride and in gadolinium oxide and sulphate are shown in figs. 28–32.

The spectrum of terbum extends a considerable distance into the ultraviolet. In addition to the bands shown in the diagram, the system Tb<sub>2</sub>O<sub>4</sub>—CaO shows the following —

```
5030; 4945, 4895, 4855; 4825, 4785–1730, 4670, 1610, 4620–1570; 4472; 4455; 1110, 1110; 4395; 4370; 4350; 4310, 1300, 4280, 4245; 4220, 4210; 4260; 4190; 4180; 4160; 4145; 4140; 4130; 4120; 3915; 3900; 3880; 3865–3840, 3620; 3805; 3795–3780; 3770; 3750; 3735; 3720.
```

<sup>&</sup>lt;sup>1</sup> Its pale brown colour (due to terbium) excepted.

The system  $\mathrm{Tb_2(SO_4)_3}$ — $\mathrm{Gd_2(SO_4)_3}$  shows the following additional bands — 4995–4975; 4900–4830; 4730, 4580, 4420; 4370, 4355; 4300; 4195-4065; 3955, 3935, 3850; 3810, 3765.

The system Tb<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>—CaSO<sub>4</sub> shows the following bands -

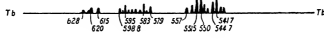
4990-4840; 4750-4675; 4590-4510; 4420; 4390; 4355;  $^{43}20$  , 4255; 4190-4080; 3865-3765;  $^{3735-3685}$ 

The system TbF3 - CaF2 shows the following bands -

5020-4890, 4890, 4825, 4705, 4650-4550, 4435, 4405; 1390, 4380-4350; 4310, 4270, 4215, 4220-4200; 4190, 4170; 4135; 3865, 3825; 3800.









 31 —Phosphorescence spectra of the rare earths diluted with gadolima (oxides prepared from oxalates)

The spectrum of terbium is of considerable interest. In 1905 Crookes announced the discovery of two new rate earth elements, incognitum and ionium, which he characterised by their ultra-violet phosphorescence spectra. The maxima were  $\lambda\lambda3760,\,3819,\,3852$  for "meognitum" and  $\lambda\lambda4351,\,4374,\,4123$  for "nomium" Reference to the figures given above will show that the "meognitum" and "nomium" spectra form part of the terbium spectrum in the system  $Tb_2(SO_4)_3$ — $Gd_2(SO_4)_4$ . The spectra assigned by Crookes to the "meta-clements"  $G_5$ ,  $G_8$ , and  $G_8$  also form parts of this same spectrum

The independent variations in the intensities of the different bands in the terbium spectrum with the dilution are exhibited in a most striking manner.

Dysprosium — The phosphorescence due to this element is pure yellow or greenish-yellow in colour. The spectra observed in calcium oxide, sulphate, and fluoride and in gadolinium oxide and sulphate as diluents are shown in figs 28-32.

Dysprosium is the element responsible for Crookes's citron band. The

<sup>1</sup> Crookes, Chem. News, 1905 92, 273

<sup>2</sup> Not to be confused with the radioactive element of that name,

correspondence of the band at  $\lambda5848~\mathrm{m}$  the system  $\mathrm{Dy}_2\mathrm{O}_3$ —CaO with the citron band  $\lambda5740~\mathrm{has},$  however, been denied by Crookes  $^1$ . The difference in wave-length is due to the fact that in his experiments the diluent was not lime but yttimm sulphate. It will be seen from the diagram (fig. 32) that when the diluent is gadolimin sulphate, the dysprosium spectrum reduces to an intense band at  $\lambda5740$  Moreover, it is found that the addition of a trace of dyspressa to a sample of yttria so pure as scarcely to show the cition band causes that band to stand out with great brilliance. The spectrum of dysprosium in yttna as diluent comprises the following bands (yellow phosphorescence) -

 $5875,\ 5850;\ 5830;\ 5790.\ 5757;\ 5740;\ 5730;\ 5705;\ 5670,\ 5610;\\ 5587,\ 5540,\ 5495;\ 5470,\ 5375,\ 5360$ 

Erbium - The phosphorescences due to erbium are either green, greenish-

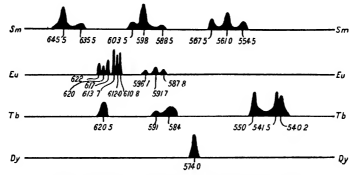


Fig. 32.—Phosphorescence spectra or rare earth sulphates diluted with gadoliuum sulphate.

blue, or blue. The spectra observed with calcium oxide, sulphate, and fluoride as diluents are shown in figs. 28, 29, and 30 <sup>2</sup>

The following bands are exhibited by erbia diluted in yttim .—

**6610**; **6540**; 5730, 5680, 5642; 5615; 5568, 5535; 5510, 5500; 5490; 5475, 5460, 5390; 5340, 5325, 5290, 5270, 5263; 5255, 5235; 5225; 5210, 5200.

Ternary Systems.- In ternary systems composed of two phosphorogens and one diluent the order of decreasing sensitiveness of the four most active phosphorogens is dysprosium, terbium, entopium, and samarium, except in oxide systems. In the latter case, the order of decreasing sensitiveness is terbum, curopum, dysprosium, sanarum, prascodymium, and neodymium. As has been already mentioned, when two phosphorogens are simultaneously present the most sensitive spectrum is slightly, and the other considerably enfeebled.

The preceding considerations indicate that when the rare earths are

Chookes, them News, 1908, 94, 79
 The orbia used contained 30 per cent. of yttim, and the Er<sub>2</sub>O<sub>3</sub>—CaO system contained 1.5 per cent. of the erbia-yttima mixture.

present to a minute extent in a mineral and the latter is examined by the method of Crookes (p. 295) for its phosphore-scence spectrum, the bands most readily observed will be the citron band of dysprosium, the orange bands of samarium, the green bands of terbium, and the "anomalous band" of europium. That the last named often fails to make its appearance is due to the extreme railty of curopium, which is only one fiftieth as plentiful as

dysprosium

Utility of Phosphorescence Spectra in Analysis.— For analytical purposes it is best to dilute a small portion of the rare earth maxime to be examined with pure lime so that the mixture contains 1 per cent of rare earth. A pure rare earth that acts only as a diluent then shows practically no phosphorescence, while a pure rare earth phosphorogen phosphoresces with its maximum power, with a rare earth mixture, on the other hand, each phosphorogen gives rise to its own phosphorescence spectrum, but not at its maximum brilliance. The phosphorogens present may be recognised by the positions of their bands and reference to fig 28. According to Urbain, the observations may be made with the same degree of certainty as may observations of are or spark spectra, with the advantage that, unlike these latter spectra, "parasite" rays (e.g. ar lines and lines of gold or platnum in the spark and carbon bands in the are) are absent

#### LINE SPECIFIC

Arc and Spark Spectra.—The line spectra of the elements as usually observed may be divided into arc and grait spectra according to the nature of the experimental procedure followed in producing them

The arc spectra of the rare earth elements are most simply produced by introducing small quantities of their oxides into the positive crater of a curbon arc light and focusing the light upon the slit of the spectroscope. Since the arc spectra are very rich in lines, the most important of which he in the ultra violet region, the spectra must be produced by means of air instrument of high dispersion, the optical parts must be made of quartz, and the spectra must be photographed. Lines and bands due to the carbon are itself are eliminated by taking, on the same plate, a comparison photograph of the spectrum of the arc, the wavelengths of the lines are found by taking another comparison photograph of a known spectrum, usually the arc spectrum of non

The spack spectra of the rare earth elements are most convemently obtained by the following method of illimination electric sparks from an induction coil or transformer are passed between a platinum, gold, or graphite electrode and a solution of the chloride or intrate of a rare earth metal, the solution being made negative to the other electrode. Here, again, the spectra should be photographed, since the most important lines he in the ultra-violet region.

Numerous convenient forms of apparatus have been destribed, and one is shown in fig.  $33^{-1}$ . A piece of glass tubing about six inches long and with a capillary bore is bent into a U-tube, as shown, and a small funnel of about 2 c.e. capacity blown at each end. A gold wire is passed through the capillary tube and made to project a little above the rim\*of the lower funnel, over its

For others, see Baly, Spectroscopy (Longmans, 2nd ed., 1912), Crookes, Proc. Roy. So., 1903, A, 72, 295, Chem. News, 1903, 88, 202.
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projecting end a short piece of thin-walled capillary tubing is slipped so that the end of the wire is level with the top of the tubing. The salt solution

ascends the capillary and keeps the end of the gold wire well wetted Another gold wire serves as the external, positive electrode.

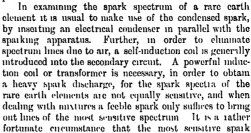


Fig 38 - Apparatus for production of spaik spectra

spectra are those of the rare earth elements that do not give rise to absorption spectra The sensitiveness of the yttrium spectrum, for instance, is very great.

When the spark spectrum of an aqueons solution of a salt is examined, it is found that as the solution is gradually diluted the lines weaken and die out at different rates. Given the same conditions, however, it was found by Hartley that the same lines always disappear at the same dilution. Hartley's work has been continued by de Gramont 2 and by Pollok and Leonard, 3 who have given the name of ultimate or residual lines to those few lines that persist the longest when the solution is diluted. It will be obvious that when the relative degrees of persistency of the various lines in the spark spectrum of an element have been determined, the observations may be advantageously employed for purposes of quantitative analysis. The results are conveniently represented in the following manner, due to Hartley lines seen in a 0 001 per cent solution of the metal are denoted by ω, those seen in a 0.01 per cent, but not m a 0.001 per cent solution by  $\psi$ , those seen in a 0.1 per cent but not in a 0.01 per cent solution by  $\chi$ , those seen in a 1.0 per cent, but not in a 0.1 per cent, solution by  $\phi$ , those only seen in solutions more concentrated than 10 per cent. by  $\sigma$ , and those seen in the metallic spark but not given by an aqueous salt solution by  $\tau$ 

In the case of the rare earth elements, the necessary measurements have only yet been effected for cerum, lanthanum, and yttimun 1

With regard to the ultimate lines in the spectrum of an element it is important to note that they are not necessarily the lines towards which a photographic plate is most sensitive (i.e. the most intense lines). The data for cerium in the accompanying table of wave-lengths will serve to illustrate this point. In a number of cases, according to de Gramont, the ultimate lines of an element are independent of the method by which the

Haitley, Phil. Trans, 1884, 175, 325.
 A de Giamont, Compt. rend., 1907, 144, 1101, 145, 231, 1908, 146, 1260; 147, 239, 807; 1909, 149, 263, 1910, 150, 1235, 151, 308, 1914, 159, 5; Ann. Chim. Phys., 1909,

spectrum is produced (flame, spark, arc, etc.). When, as often happens, this is not the case, it is nevertheless found that the ultimate lines are always lines of great persistency in the condensed spark spectrum

A number of regularities in the line spectra of some of the rare earth elements has been observed, but no definite series of lines have yet been

The accompanying table gives the wave-lengths of the most intense, and, where known, the most persistent lines in the arc and spark spectra of the rare earth elements.

### CHIEF ARC AND SPARK LINES IN THE SPECTRA OF THE RARE EARTH ELEMENTS \*

| Spark | Arc               | Spark      | Arc.     | Spark.    | Λιc     |
|-------|-------------------|------------|----------|-----------|---------|
|       |                   |            | 2949 5   | 1150 09   | 4352 33 |
| Crli  | II M <sup>1</sup> |            | 3080 7 3 | 4165 75 3 | 4356 0  |
|       | 2159 1            |            | 3118 6 2 | 4181 25 3 | 4460 4  |
|       | 2169 3            |            | 3171 /   | 4186 71   | 1479 5  |
|       | 2481 6            |            | 3197 92  | 4290 88   | 4157 0  |
|       | 2536.9            | 2.0        | 3326 0   | 1391-81   | 1527 5  |
|       | 2677 7            |            | 3 :91 5  | 1160 19   | 4528 6  |
|       | 2055 22           | 1          | 3665 6   | 3527:527  | 4539 9  |
|       | 27.29 1           |            |          | 4562 52   | 4562 5  |
| •     | 277 9             | Ciri       |          | 4572 16   | 1572 4  |
|       | 2765 8            |            |          | 1591 12   | 4594 1  |
|       | 25313             | 260 - 72 ' | 3952.77  | 4628 33 i | 4624 3  |
|       | 2537 3            | 30.666     | 1119 99  | 5079 5    | 5353 7  |
|       | 2845.2            | 5801.71    | 4150-11  | 5511 23   | 5512.2  |
|       | 2870 2            | 5930 (93   | 41~6 78  |           | 6272 2  |
|       | 2855 1            | 4012.58    | 1222.78  |           | 6371 3  |
|       | 2003.9            | 1133 98    | 1295 85  |           |         |
|       | 2931.7            | 1137 78    | 4337 96  |           |         |

Only the range \$2450 to \$3700 has been examined.

Pollok, Set Proc Roy Itabl. Sr., 1912, 13, 202–253
 See Hicks, Phil. Trans., 1912, A, 212, 33, 1913, A, 213, 323 (Sc., Y., La, Gd., Yh).
 Popov, Ann. Physik, 1914, [iv], 45, 147 (Yv., Panlson, dvd., 1914, [iv], 45, 419 (Sc., Y),
 1203 (La), Physikal Zatish. 1914, 15, 892 (Y., Sc., Lu), 1915, 16, 7 (Gd.), 352 (Tm)

The most intense arclines of celtium

Begree of persistency equal to x

<sup>&</sup>quot;With the exception of the lines of collisms, the lines given in these tables in ordinary and in clarendon types represent the "haupthmen" or most interestines, and are taken from Exner and Haschek, Die Sychten der Elemente ber normalem Priek Leipzig and Viennis, 1911)—Lines given in italies are among the most persistent but not among the most intense lines. Where they are known, the relative persistent but not among the most intense lines. Where they are known, the relative persistent but not among the most intense lines. Where they are known, the relative persistent of the spark lines are indicated in Hartley's notation (p. 306)—Lines of intensity 50 or over on Exner and Haschek's scale of intensities are given in heavy type.

The wave-lengths given by Exner and Haschek are correct to within one or two units in the second deermal place. Bearing this in mind it is easy to pick out the "haupthmen" of a rare carth element that we common to the are and spark spartia.

Spark Spectra.—The following is not a complete bibbography, but includes practically all the modern references to the spark spactra of the rare earths.—

Cerium.—Brauner, Monatsh., 1882, 3, 1, Lohse, Publ. Astrophys. Observ. Potsdam, 1902, 12, 109, Pollok and Leonard, Sci. Proc. Roy. Dubl. Soc., 1908, 11, 267, Exner and Haschek, Sitzungsber, K. Akad. Wiss. Wien, 1899, 108, 11a, 1071, and opus cit.

# CHIEF ARC AND SPARK LINES, ETC .- continued

| Spark.      | A1c.     | Spark,          | Aıc.               | Spark.               | A1c.    |
|-------------|----------|-----------------|--------------------|----------------------|---------|
| Dysphosium. |          | Dysprosium. 372 |                    | 4129'90 <sup>2</sup> | 3972.10 |
| 3154 50     | 3524.14  |                 | 3830 69            | 4205 20 3            | 4129'90 |
| 3521 20     | 3524 18  |                 | 3895.40            | 4435'74              | 4205'20 |
| 3531 881    | 3531 .87 |                 | 3902.95            | 4522 75              | 4435'7  |
| 3550:351    | 3536 20  |                 | 3906 17            | 4594 28 3            | 4522 7  |
| 3600 19     | 3538 66  |                 | 3932 48            | 6645 41              | 4594.5  |
| 3645.55     | 3538 70  |                 | 3938.79            |                      | 4627.47 |
| 3076 71     | 3550 37  |                 | 3973 26            |                      | 4662'10 |
| 3694 99     | 3941 83  |                 | 3973 78            |                      | 6645.44 |
| 3698 30     | 3948:56  |                 | 3974 89<br>4008 12 |                      |         |
| 3806 14     | 4000 61  |                 | 4020:69            |                      |         |
| 3598 70     | 4000 69  |                 | 4059 98            | GADOLI               | NIUM.   |
| 3911 821    | 1046 16  | 1               | 4087 80            | 2628 22 4            | 3082:18 |
| 3978 721    | 4078 11  |                 | 4143 11            | 2901 84              | 3100 66 |
| 1000 61     | 1103 50  |                 | 4151.29            | 2955 62              | 3122 62 |
| 4078.15     | 4168 15  |                 | 4119.78            | 3350 66              | 3515 9  |
|             | 4187 00  | i               | 4563 45            | 3362 41              | 35 19 5 |
|             | 4211 89  |                 | 4631.10            | 3122 65              | 3585 12 |
| ł           | 4211 89  | i i             | 1675 77            | 3515 954             | 3646 36 |
|             | 4221 30  |                 | 4759 83            | 3549 51 4            | 3671:39 |
|             | 4589 53  |                 | 5827 01            | 3585 10              | 3719 63 |
|             | 4957 59  |                 | 6221 22            | 3646 32              | 3713 68 |
| •           |          |                 |                    | 3661 76              | 3768'60 |
| Err         | IIIM     |                 |                    | 3712 89              | 3796 65 |
|             | -        | EUROI           |                    | 3719.62              | 3811 18 |
| 3372 91     | 3312.60  |                 | TU M               | 3713 60              | 8852 68 |
| 3199 29     | 3372.92  | 3725 08         | 3688 57            | 3768 51              | 3916.70 |
| 3692 80     | 3355 23  | 3810.81 3       | 3725 10            | 3782 51              | 4037.49 |
| 3906 51     | 3199 28  | 3907 30         | 3819 80            | 3791, 58             | 4050 08 |
| 4419 79     | 8616 75  | 3930 652        | 3907 28            | 4130 55              | 4063 62 |
| 1675 80     | 3692 85  | 3972'16"        | 3930 66            | 4184 18              | 4071 51 |

<sup>1</sup> Dem up w'e element  $\Delta$ . The other lines ascribed to  $\Delta$  by the same authority are \$\text{\$\lambda\$}(212.6, 4195.5, 4187.3, and 3595.0. they are presumably very persistent dysprosium lines.

1 Demargay's element  $\Sigma$ 1 Lecoq de Borsbaudian's element  $Z_c$ . The other lines ascribed to  $Z_c$  by the same authority are \$\text{\$\lambda\$}(627.0)\$ and \$4602.00\$, they are presumably very persistent curopium lines.

4 An ultimate line, according to Demargay.

Dysprosium.—Lecoq de Borsbaudian, Compt. rend., 1886, 102, 153, 1003, 1005, Demaicay, it id., 1900, 131, 387, Urbain, ibid., 1908, 146, 922, Exact and Haschek, opus cit

Erhum.—Thalen, Compt. rend., 1880, 91, 326; Orteors, K. Vet Akad Forhundt, 1881, 6, 13; Exner and Haschek, Sitzungsher K Akad. Wiss Wien, 1899, 108, 11a,

<sup>1881, 0, 13,</sup> Evnet and Haschek, Sitzingsher K Akad. Wiss Wien, 1899, 108, 11a, 1123, and opus cit

Europium.—Demateay, Compt. rend. 1896, 122, 728; 1900, 130, 1019, 1469; 1901, 132, 1484; Evnet and Haschek, Sitzingsher K. Akad. Wiss. Wien, 1902, 111, 11a, 42, and opus cit; Crookes, Five Roy Soc, 1905, 74, 550.

Gadolinium.—Leven de Borshaudhan, Compt. rend., 1886, 102, 902, 1889, 108, 165; 1890, 111, 472, Demateay, ibid., 1896, 122, 728, 1900, 131, 343, 995, Exnet and Haschek, Sitzingsher, K. Akad. Wiss. Wien, 1900, 109, 11a, 103; 1901, 110, 11a, 557, and opus cit

and opins of
Holmium.—Evner and Haschek, Sitzungsber, K. Akail. Wiss, Wien, 1901, 110, Ha,
557, 1910, 119, Ha, 771, and opins et.
Lanthanum.—Brauner, Monatsh., 1882, 3, 1, Bettendorf, Annalen, 1890, 256, 159;
Exner and Haschek, opus et., and Sitzungsber, K. Akail. Wiss, Wien, 1899, 108, Ha,

### CHIEF ARC AND SPARK LINES, ETC .- continued.

| Spark                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | Aie                                                                                                                                                                                   | Spark.                                                                                                                                                                                           | Arc                                                                       | Spark                                                                                                                 | Are                                                                                                                                                                                                                                                      |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Horal State of State | 4073 99 4085 73 1098 80 4130 59 1181118 4251 90 4262 21 4325 83 1327 29 1342 45 6114 26 MI W. 3399 12 3425 49 3425 49 3428 27 3156:15 3471 10 3141 98 3518 73 3798 92 3718 32 3757:41 | 2297 904<br>2379 494<br>2376 891<br>2576 891<br>2508 464<br>3171 794<br>3517 26<br>3759 33<br>3791 02<br>3794 99<br>388 661<br>1041 86<br>4043 18<br>4085 90<br>4383 974<br>4522 644<br>4907 704 | 3849 27<br>3988 69<br>4238 55<br>4333 98<br>6250 11<br>6252 52<br>6394 16 | 2895 00<br>2911 59<br>2963 11<br>3058 08<br>3077 75<br>3254 41<br>3397 16<br>3172 62<br>3.07 56<br>3554 58<br>4181 11 | 3198 27<br>3254 46<br>3281 89<br>3312 30<br>3359 74<br>3376 69<br>3397 21<br>3472 65<br>3508 55<br>3508 55<br>3508 55<br>3568 90<br>3624 10<br>3636 41<br>3876 80<br>4124 87<br>4184 40<br>4518 74<br>5476 88<br>5683 92<br>5684 32<br>6646 40<br>663 40 |
| 3515 74<br>3556:95                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 3889 10<br>3891 17                                                                                                                                                                    | 5588 54 1                                                                                                                                                                                        |                                                                           | Nioby                                                                                                                 | MILM                                                                                                                                                                                                                                                     |
| 3598 92<br>3748:35<br>3757:40<br>3796 91<br>3810 91<br>3891 18                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 1 4015 58<br>4251 59                                                                                                                                                                  | Le (1<br>(case o)<br>2603 41<br>2615 52<br>2772 70                                                                                                                                               | cium<br>oreum).<br>2615:50<br>2911-53<br>3077-75                          | 4012 12<br>4031 93<br>1061 26<br>4156 30<br>4177 50<br>4303 77                                                        | 8863 52<br>3951 32<br>4061 27<br>4156 50<br>4247 54<br>1282 67                                                                                                                                                                                           |

<sup>&</sup>lt;sup>1</sup> Degree of persistency equal to  $\chi$ <sup>2</sup> Degree of persistency equal to  $\psi$ <sup>3</sup> Cleve's "clement" Di $\beta$ .

<sup>1071.</sup> Edet and Valenta, thel., 1909, 118, 11a, 511, Lehse, Publ. Astrophys. Observ. Potsdam, 1902, 12, 109; Pollok and Leonard, See Proc. Roy. Inth. Soc., 1908, 11, 257.

Lutecium—Uthain, Compt. read., 1907, 145, 759, 1908, 164, 400; Auci. von Welsbach, Stexingsber. K. Akad. Wiss. Wica, 1907, 116, 115, 1125, 1909, 118, 116, 507; Exner and Haschek, thel., 1899, 108, 11a, 1123, and opins ed., Crookes, yede intra.

Neodymium.—Auci. von Welsbach, Monatsh., 1885, 6, 177; Sidzungsber. K. Akad. Wiss. Wica, 1885, 92, 11, 317, 1903, 112, 11a, 1037; Exner and Haschek, thel., 1899, 108, 11a, 1021, and ones ed.

<sup>Wiss, Wien, 1885, 92, II 317, 1903, 112, IIa, 1037, Exnet and Haschek, ibid., 1899, 108, IIa, 1071, and opics cit.
Prascodymum.—Auet von Welsbach, ibid., Exnet and Haschek, ibid. at Samarium.—Bettendorf, Annalen, 1891, 263, 164, Demaryy, Compt. rend., 1900, 131, 995; Exnet and Haschek, Stitumpher K. Ikad. Wiss. Wien, 1900, 109, IIa, 103, 1901, 110, IIa, 577, and opits cit.
Terbium.—Demaryay, Compt. rend., 1900, 131, 387, Utbain, ibid., 1906, 142, 957;
J. Chem. phys., 1906, 4, 349, Exnet and Haschek, opits cit.
Thulium.—Thalen, Compt. rend., 1880, 91, 376, Exnet and Haschek, opits cit., and Stitumysber. K. Akad. Wiss. Wien, 1901, 110, IIa, 964; 1910, 110, IIa, 771. Auet von Welsbach, ibid., 1911, 119, IIa, 193; Mondish., 1911, 32, 373. Nettch. anory. Chem., 1911, 71, 439, Crookes, Proc. Roy. Soc., 1906, A, 78, 154; Utbain, Compt. rend., 1907, 145, 759.
Yttrum.—Hattey, Trans. Chem. Soc., 1882, 41, 210, Exnet and Haschek, Nitsungaber. K. Akad. Wiss. Wien, 1899, 108, IIa, 1123, and opus.cit.; Pollok and Leonard, Scit. Proc. Roy. Dubl. Soc., 1908, 11, 257.</sup> 

# CHIEF ARC AND SPARK LINES, ETC .- continued.

| Spark.   | Arc.      | Spark     | Aro.               | Spark.               | Arc.               |
|----------|-----------|-----------|--------------------|----------------------|--------------------|
| 4446.54  | 4303.78   | 4409.01   | 4429:38            |                      | 4544-11            |
| 4451 '68 | 4325.87   | 4429.41 . | 4496 60            |                      | 4566 3             |
| 4463.12  | 4875-11   | 4496 63   | 4510.32            |                      | 4577 88            |
|          | 4385.81   | 4510.33   |                    |                      | 4642.4             |
|          | 4400.96   |           |                    |                      | 4674 7             |
|          | 4446.21   | SAMA      | RIUM.              | '                    | 20/2/              |
|          | 4451.71   | 4319.12   | 3789:30            | TERBI                | IIM. 1             |
|          | 4463.09   | 4391.05   | 4152 38            | 2659.01              | 3324.58            |
|          | 4920.84   | 4424.52   | 4203.18            | 2891.40              |                    |
|          | 5293:35   | 4434.50   | 4225.48            | 2909:35              | 3509 34            |
|          | 5319 98   | 4467:49   | 4229 83            |                      | 3581 86            |
|          | 5594.58   | 110/ 15   | 4236.88            | 2913 39<br>3509 32 2 | 3561.90            |
|          | 5620.75   |           | 4256 54            |                      | 3568-69            |
|          | - 6310.69 |           | 4319-12            | 3561 90 <sup>2</sup> | 3600 60            |
|          | 6341.69   |           | 4329.21            | 3676.522             | 3628 35            |
|          | 6385:32   |           | 4834.32            | 8703 00°             | 3650.60            |
|          |           |           | 4347.95            | 3747:48              | 8659.02            |
| Praseci  | DYMIUM.   |           | 4391.03            | 3848 90              | 3704.10            |
| 8877 38  | 4008 90   |           | 4420.72            | 3874 32              | 3711-91            |
| 4100.89  | 4100.91   |           | 4421.32            | 3925.60              | 3848 90            |
| 4118.63  | 4118.70   |           | 4424.55            | 8939.71              | 3874 38            |
| 4143.28  | 4148:33   |           |                    | 3977.00              | 3899:34            |
| 4179.60  | 4179 60   |           | 4434·07<br>4434 52 | 8892.04              | 3925.60            |
| 4189.70  | 4189.70   |           |                    | 4144.60              | 3939 75            |
| 4206.81  | 4206.88   |           | 4452-92            | 4278 67              | 3977 01            |
| 4223.20  | 4223.18   |           | 4454.84            | ' '                  | 3982.07            |
| 4225.54  | 4225 50   |           | 4458.70            |                      | 4005 70            |
| 4241.28  | 4225 50   |           | 4467.50            |                      | 4012 99            |
| 4805 99  | 4305.99   |           | 4519·80<br>4524 08 |                      | 4278 70<br>4752:69 |

According to Eberhard, the most persistent lines in the arc spectrum of terbium are λλ3523.82, 3676.52, 3703.00, 3704.10, 4005.70, 4278.70.
 Demarçay's element Γ. The other lines ascribed to Γ by that authority arc λλ3523.79, .3540.42, 3568.68, and 3704.08, which are presumably very persistent terbium lines.

Ytterbium (neoytterbium).—Urbain, Compt rend., 1907, 145, 759; 1908, 146, 406; Auer von Welsbach, Sitzungsber. K. Akad. Wiss. Wien, 1907, 116, 116, 1425, 1909, 118, 116, 507; Exner and Haschek, opus cit. Crookes, loc. cit.

Arc Spectral.—The following is not a complete bibliography, but includes practically all the modern references to the arc spectra of the rare earths:—

General.—Exner and Haschek, opus cit. (Ce, Dy, Er, Eu, Gd, Ho, La, Lu, Nd, Pr, Sm; Tb, Tm, Y, Yb); Eder and Valents, Sitzungsber. K. Akad. Wiss. Wien, 1910, 110, 114, 3 (Lu, Yb), 9 (Dy, Tb, Er, Gd, Ho, Y, La, Sm, Eu), 103 (Tm), 519 (Lu, Ce, La, Nd, Pr, Y).

Celtium.—Urbain, Compt. rend., 1911, 152, 141.

Cerium.—Bakowski, Zeitsch. wiss. Photochem., 1908, 6, 73; Cooper, Astrophys. J., 1909, 29, 329; Eder, Chem. Zentr., 1915, i. 1046.

Dysprosium.—Eberhard, Zeitsch. wiss. Photochem., 1906, 4, 137; Astrophys. J., 1906, 34, 309; Publ. Astrophys. Observ. Potsdam, 1909, 20, 3.

Etropium.—Exner and Haschek, Sitzungsber. K. Akad. Wiss. Wien, 1902, 111, IIa, 342, and opus cit.

Gadolinium.—Exner and Haschek, Sitzungsber. K. Akad. Wiss. Wien, 1902, 111, IIa, 342, and opus cit.

Gadolinium.—Eberhard, Publ. Astrophys. Observ. Potsdam, 1909, 20, 3.

Lanthanum.—Rowland and Harrison, Astrophys. J., 1898, 7, 373; Kellner, Inaug, Miscolinium. Photochem., 1905, 3, 395; Humphreys, Misrophys. J., 1907, 26, 18; Eder, Chem. Zentr., 1915, i. 1046.

# THE RARE EARTH ELEMENTS.



### CHIEF ARC AND SPARK LINES, ETC .- continued.

| Spark.            | Arc.               | Spark     | Arc     | Spark.    | Arc.      |
|-------------------|--------------------|-----------|---------|-----------|-----------|
| Тни               | IIIM.              | 2891.50   | 3289:50 | 3633.28   | 3982 79   |
| 3181.40   3181.40 |                    | 2919:49   | 8464 47 | 3664.76   | 4077.54   |
| 8184.01           | 3131 40<br>3134 00 | 8005 85   | 3694'35 | 3710'44 3 | 4102.57   |
| 3151 18           |                    | 3026 78   | 3988.10 | 3774'51 3 | 4128.50   |
|                   | 8362-78            | 3108 00   | 5556.67 | 3788 88   | 4148 03   |
| 3172 98           | 8425 27            | 3193 01   |         | 3850·51   | 4177.74   |
| 3286.96           | 3441 71            | 3289.50   |         | 3982.78   | 4302 45   |
| 3261.75           | 3453 82            | 3875.65   |         | 4177.68   | 4809.79   |
| 3362.75           | 3462 87            | 3478-99   |         | 4309.81   | 4348 93   |
| 8425 25           | 3608-92            | 3694°35°2 |         | 4375'11 4 | 4375'12   |
| 3462.88           | 3700.41            | 3988 202  |         | 4855.06   | 4883 89   |
| 8700 40           | 3701 54            | 4726:24   |         | 4883.053  | 6191.91   |
| <b>3</b> 701 52   | 8718 07            | 4786 82   |         | 4900.353  | 6435-27   |
| 3761 50           | 3734 29            |           |         | 5087·613  |           |
| 3762.05           | 8744-22            |           |         | 5663.17   |           |
| 3795.50           | 3761 49            |           |         |           |           |
| 3848-14           | 3762 09            | YTTR      | IUM.    | }         |           |
| 4522.76           | 8795.90            | 2367 30   | 3216.83 | DEMARÇ    | 1 T'O O   |
|                   | 3848.13            | 2414 783  | 8242.42 | DEM ARI,  | A Y B 12. |
|                   | 4094.33            | 2422 303  | 3328.02 | 3930 9    |           |
|                   | 4105 99            | 2817 143  | 3600.92 | 3967 9    |           |
|                   | 4187.79            | 2946 15 3 | 3611:20 |           |           |
|                   | 4481.44            | 3216 84 3 | 8621.10 |           |           |
|                   |                    | 8242 40 4 | 8633.28 |           | -1        |
| YTTEI             | BIUM <sup>1</sup>  | 3328.03   | 8664.78 | DEMARÇ    | A T'S 0.  |
|                   | um or alde-        | 8549 12   | 8710.47 | 3906-5    |           |
| baran             | ium).              | 3600.003  | 8774.52 | 4008-2    |           |
| 2803.55           | 3031.26            | 3602.12   | 3788.88 |           | i         |
| 2818 39           | 3107.99            | 3611-193  | 3950.52 |           |           |

<sup>&</sup>lt;sup>1</sup> The data given in the table for ytterbium are taken from Exner and Haschek, opus ett. At the present time (1916) the only spectrum measurements that have been taken with pure ytterbia are the measurements of the aic lines effected by Blumenfeld and Uibain (Compt. rend., 1914, 159, 401). The most intense are lines between  $\lambda = 2300$  and  $\lambda = 3500$  are as follows, according to these observers:—

2390'8, 2398'1, 2464'5, 2512'1, 2538'7, 2596'4, 2617'1, 2642'0, 2644'3, 2653'9, 2665'1, 2672'8, 2684'8, 2710'7, 2718'5, 2732'9, 2750'6, 2771'5, 2776'4, 2784'8, 2814'6, 2821'3, 2825'1, 2831'1, 2847'4, 2851'2, 2861'5, 2867'3, 2891'5, 2919'5, 2946'0, 2970'7, 2984'3, 2995'0, 3005'9, 3031'3, 3065'2, 3089'2, 3107'9, 3193'0, 3201'2, 3226'0, 3261'6, 3289'4, 3337'3, 3343'0, 3476'4, 3479'0, and 3496'0.

- $^2$  The most persistent lines of ytterbium, according to de Gramont.  $^3$  Degree of persistency equal to  $\chi.$   $^4$  Degree of persistency equal to  $\psi.$

Lutecium.—Exner and Haschek, opus cit; Eder and Valenta, loc. cit.
Neodymium.—Bertram, Zeitsch. wiss. Photochem., 1906, 4, 16.
Praseodymium.—Bertram, loc. cit.
Samarium.—Rutten, Zeitsch. wiss. Photochem., 1905, 3, 181.
Terbium.—Eberhard, ibid., 1906, 4, 137; Sitzungsber. K. Akad. Wiss. Berlin, 1966 384.
 Thulium.—Exner and Haschek, opus cit.; Urbain and Blumenfeld, Compt. rend.,

914, 159, 323, 401. Yttrium. - Kayser, Abhandl. Berlin Akad., 1903; Eberhard, Zeitsch. wiss. Photochem.

The X-Ray or High Frequency Spectra.—X-rays are emitted by an element when the element or one of its compounds is bombarded with cathode rays. The X-rays are ether-waves of very small wave-length, i.e. of very high frequency, and the spectrum of the X rays emitted by an element may be examined by a process analogous to the diffraction grating method for the examination of ordinary spectra. For a description of the process the reador is, however, referred elsewhere, as it is beyond the scope of this book 1

The X-ray spectra of the elements have been examined by Moseley and others 2 The spectra are extremely simple, consisting as a rule of four or five lines at the most. When, therefore, the necessary apparatus is available, the study of these spectra should be preferable to that of the are or spark spectra, which are extremely complex. Preliminary wave-length measurements have been made by Moseley for the following rare earth elements: lanthanum, cerum, prascodymium, neodymium, samarium, europium, gadolimum, dysprosium, yttrum, and erbinn.

The X-ray spectra of the rare earth elements throw a good deal of light on the question of the homogeneity of these substances. Further discussion of these spectra is therefore given in the following chapter (see p. 363)

### APPLICATIONS OF THE RARE EARTHS!

According to a statement by Stein 5 in 1913, the incaudescent mantle industry consumes about 3300 tons of monazite sand per annum, 300 tons of thorum intrate being produced. Very large quantities of earths of the cerum group are obtained as by-products, in fact, about 1000 tons of cera per annum are obtained, only 3 tons of which are required by the mantle industry. The mixed rare earth ovalates or carbonates left as residues are usually known as crude cerium oxalate or carbonate, and their profitable utilisation in chemical industry is a problem that has at present received no really satisfactory solution, although many applications of them have been suggested A number of these may be briefly discussed.

Gas Mantles and Electric Filaments,-An ordinary meandescent gas mantle contains about 0 8-1 0 per cent of cena, CcO, and a few tenths of a per cent. of glucma, GlO, and alumma, Al<sub>2</sub>O<sub>3</sub>, the remander being composed of thoria, ThO<sub>2</sub>. In mantles designed for use with high-pressure gas the percentage of cerm is increased and may reach the value 2 8 per cent.

As has been already mentioned, the annual consumption of ceria in the mantle industry is very small To an even smaller extent "didymia" is also employed. When mantles are required to be marked they are stamped, just before the burning-off process, with a solution of "didymium" intrate mark then appears on the finished mantle as a pale brown, permanent coloration.

See Kaye, X. Raus (Longmans & Co., 1914), and the papers cited in the next footnote
 Moseley, Phil. Mag, 1913, [vi], 26, 1024, 1914, [vi], 27, 703, Malmer, ibid., 1914, [vi], 28, 787, Rydberg, ibid., 1914, [vi], 28, 144, de Broghe, Compt. rend., 1914, 159,

There is a small demand for yttim in connection with the manufacture of Nernst lamps - In these lamps illumination is produced by passing an electric current through a hlament composed of a mixture of refractory oxides, of which yttra, thoria, and zircoma are the most sintable for the purpose 1. As a source of yttria the gadolinite deposits found at Barringer Hill, Llano Co., Texas, are utilised

Alloys. The production of mischmetall, an alloy of the metals of the cerum group, has been mentioned earlier in this chapter (p. 230). Owing to the great stability of the oxides of the rare earth elements, these elements and certain of their alloys form valuable reducing agents. Mischmetall and the cernin magnesium alloys have been especially mentioned in this connection  $^{2}\,$ 

The ignition temperature of cerium in an is very low. Firther, numerous alloys rich in cerruin are hard and brittle, so that when struck with hard steel small particles of the alloys are detached and at the same time heated sufficiently to cause them to ignite - These alloys are usually called pyrophoric alloys, the preparation and uses of many of them are protected by patents. They may be prepared either from cermm or from mischmetall, the alloys containing 30 per cent of non (mekel or cobalt) have been particularly specified by Aner von Welsbach, and alloys with tin, lead, zinc, cadminn, silicon, titanium, etc., have also been patented 1. The alloys are used in the manufacture of automatic eigarette-lighters, gas-lighters, etc. It has also been proposed 5 to utilise them as are lamp electrodes, either alone or as the cores of carbons. According to Stern,6 200 tons of ceria are utilised annually in the production of pyrophoric alloys?

According to Husch, cerum-magnesium alloys rich in cerum form excellent flashlight powders?

It has been claimed that the introduction of 0.2 per cent of cerium into aluminium 10 greatly improves the mechanical properties of the metal 11

Arc-lamp Electrodes .- The incorporation of cernin compounds, particularly cerimin fluoride, into the carbons used as arc-lamp electrodes is the subject of various patents 12 lt is claimed that a very intense light is thereby obtained and the arc burns evenly and quietly. One impregnating mixture proposed for the purpose contains 60 per cent, of the mixed mulybdates and tungstates of the cerum group, 20 per cent of the corresponding iron salts, and 20 per cent of the fluorides of the ceruin group

According to Stern, 13 300 tons of cerm are consumed annually in the form of fluoride for impregnating arc-light carbons

See Eng Pat., 1898, No. 6135
 See pp. 231, 379, and Mathinann and Beck, tunalen, 1904, 331, 46. Huseb, J. Ind. Eng Chem., 1911, 3, 880, Trans. Timer. Electrochem. Soc., 1911, 29, 57., Werss and Archel, Annalen, 1905, 337, 370. Kellermann, Die Certimetalle und ihre propheren. Legierungen (Knapp, Halle, 1912). Escales, D.R.P., 145,820 (1903)
 Auer von Welsbach, D.R.P., 154,807 (1903). Eng. Pat., 16,853 (1903)
 See, e.g., Eng. Pat., 27,341 (1908). Fr. Pat., 439,058 (1912)
 See Eng. Pat., 1909, Nos. 8150 and 8151.
 Stein, Lie, etc.

<sup>6</sup> Stern, luc. cit.

<sup>7</sup> For further information on these alloys, see Kellermann, opus cit, Olsen, Polytech. Eng , 1914, 14, 24 8 Husch, loc est

Cerium Glass.—The incorporation of ceria into an or linary colourless soda glass results in the production of a glass that has at the most a faint yellow tinge. It is very transparent to luminous rays, but cuts off practically all the ultra-violet and about 30 per cent of the heat rays. By introducing other special ingredients into the cerium glass, a valuable series of glasses has been prepared by Crookes. The compositions of the mixtures employed in their preparation are given in the following table (the "raw soda flux" has the composition, sand 61 00, anhydrous sodium carbonate 25 50, sodium nitrate 5 00, calcium carbonate 7 20, borax 0 75, and arsenious oxide 0 55 per cent, and the "fused soda flux" is obtained from the raw flux by fusion)—

| [                                                                            | , -         | ı     | , -   | ı    | ı     | ı    |      |       |       |       | ì     | ,     |
|------------------------------------------------------------------------------|-------------|-------|-------|------|-------|------|------|-------|-------|-------|-------|-------|
|                                                                              | 1           | 2     | 3     | 4    | 5     | 6    | 7    | 8     | 9     | 10    | 11    | 12    |
|                                                                              | -           | '     | ' -   |      | -     | ļ    |      |       |       | 1     | ·     | ļ     |
| Raw soda flux .<br>Fused soda flux                                           | 90 00       | 89 75 | 87 56 | 83 0 | 79 00 | 50.0 | 77.0 | 92 00 | 94 60 | 89.17 | 98 00 | 72 60 |
| CelsO <sub>1</sub> , 6H <sub>2</sub> O                                       | <br>  8   3 | 8 18  | l     | 17:0 | 20 50 | 13.1 | 23 0 | 6:30  |       | 10 00 | 5 00  | 24 90 |
| N1SO <sub>4</sub> 711 <sub>2</sub> O<br>CoSO <sub>4</sub> 711 <sub>2</sub> O | 0 07        |       |       |      | 0 30  |      |      |       | 0 08  | 00;   | 000   |       |
| FeSO, 711 O<br>CuSO, 5H O<br>Fe <sub>3</sub> O,                              | <br>        |       | 3 00  |      |       |      |      |       |       | .     | 1.12  | 2 10  |
| Fe <sub>2</sub> O <sub>3</sub><br>Cr <sub>2</sub> O <sub>3</sub>             | "           | 0 09  | 0 80  |      | :.    |      |      | 1 60  |       | 1 50  |       |       |
| N <sub>1</sub> O                                                             | 1 80        | 0 03  | 0 09  | •    | 0 15  | 6.6  |      | 0.00  | 0 .30 |       | 0 10  | 0.10  |
| UO <sub>3</sub>                                                              |             |       | 0 55  | •••  | 0 10  |      |      |       | 0 30  |       | 2 75  |       |

The next table gives the tints of these glasses, the percentages of heat rays absorbed and visible rays transmitted, and the wave-length beyond which the ultra-violet is absorbed:—

| No of<br>Glass                                                | Tint of Glass.                                                                                                                                                              | Heat Rays<br>Absorbed                         | Lummous Rays<br>Transmitted                  | Ultra-violet<br>Absorbed.                                                                            |
|---------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------|----------------------------------------------|------------------------------------------------------------------------------------------------------|
| 1<br>2<br>3<br>4<br>5<br>6<br>7<br>8<br>9<br>10<br>11<br>- 12 | Pale yellow Pale greenish-yellow  Almost colourless Pale neutral Pale yellow Almost colourless Pale green Dark neutral Very pale blue Pale greenish-yellow Pale blush-green | Per cent. 37 63 38 27 41 39 34 29 47 51 25 47 | Per cent 73 54 48 99 45 60 71 71 30 63 74 45 | hom<br>33013<br>3700<br>3880<br>3650<br>3700<br>3685<br>3610<br>3620<br>3650<br>3650<br>3685<br>3685 |

The preceding glasses are of considerable value for the construction of spectacles Glass 10 is fairly transparent and does not perceptibly alter the

<sup>1</sup> Crookes, Phil. Trans., 1914, A, 214, 1; Chem. News, 1914, 109, 265, 277, 289, 301.

colours of objects viewed through it; since it cuts off 51 per cent. of the heat rays, it is well adapted for making spectacles for those employed in glass works. Glasses 1, 2, 5, and 6 are good absorbers of ultra-violet rays, numbers 1, 4, 7, 8, and 11 are the most transparent to visible rays, while for glare-reducing spectacles numbers 3, 5, 9, 10, and 12 are emmently suitable.

Tanning .- It has long been known that certain aluminium, iron, and chromium salts are capable of converting hides into leather, and chrome tanning is of considerable importance. According to Garelli, salts of cerium, lanthanum, and "didymnim" have tanning properties similar to those of alummum salts, while ceric salts are particularly well adapted for the process of taining. Parenzo agrees with Garelli's views, but Either is of the contrary opinion 1

Photographic Uses.—A solution of cerie sulphate in dilute sulphuric acid has a certain limited application in photography as a reducer of over-developed negatives, its action being even and rapid. Ceric salts have also been incorporated into photographic emulsions for pigment printing. The printing paper is coated with an emulsion prepared by adding the cere salt to a colloidal solution of albumin in aqueous ammonia, sodium carbonate, or borax. The paper, which is very sensitive to light, is exposed in the usual way and developed in a suitable solvent for the albumin, e.g. aqueous ammonia, sodium carbonate, or potassium ovalate 3

Dyeing .- Cerum compounds have been at times proposed as mordants, eg. in dyeing with alizarin. They do not, however, appear to be used

Rare Earth Catalysts .-- The crude mixture of basic sulphates of the cerum group has been proposed as a suitable catalytic agent for use in the contact process for manufacturing sulphuric acid. The mixture is said to be more effective than the pure cerous compound. The application of the oxides of the rage earth elements to the same process has also been proposed proposals do not, however, appear to be of much practical value 5. Cernun mittide is said to hasten catalytically the union of mitrogen and hydrogen, and its use in the production of synthetic ammonia has been patented 6

Cerous sulphate is said to form an excellent catalyst for use in the oxidation of aniline to anilino black ?

It may also be mentioned that ceria has been proposed as a sintable substitute for platinised asbestos in Dennstedt's method for the combustion of organic compounds b

Oxidation of Organic Compounds.—Since cere salts are readily reduced to the cerous state, they may be utilised as oxidising agents. It is

<sup>1</sup> Garelli, Atti R. Accad Lincei, 1907, [v.], 16, 1.532. Collegium, 1912, p. 418. J. Soc. Chem. Ind., 1912, 31, 830. Parenzo, C. Hagiam, 1910, p. 121; J. Soc. Chem. Ind., 1910, 29, 579., Eitner, Gerber, 1911, 37, 199, 213. J. Soc. Chem. Ind., 1911, 30, 1128.

2 Lumbre, Bull. Soc. Franç. Photog., 1900, 16, 103. Lumbre Bros. and Seyewetz, Monsteir Sixent, 1901, 15, 169. See also Bird. J. Photog., 1900, 47, 184, 1901, 48, 594; and Eng. Pat., 470 (1900).

3 Galeau, F. Pat., 380, 188 (1907). Eng. Pat., 20,740 (1907).

4 See Witt, Chem. Ind., 1896, 19, 156. J. Soc. Chem. Ind., 1896, 15, 580; Matschak, 1914, 1898, 17, 150., Gueliu, 1904., 1898, 17, 577.

5 See Eng. Pat., 1385 (1901). Fr. Pat., 326, 321 (1903), and Pluddemann, Inaugural Dissectation (Berlin, 1902).

Dissertation (Berlin, 1907)

loc. ett.

8 Bekk, Ber., 1913, 46, 2571, Miss M. Reimer, J. Amer. Chem. Soc., 1915, 37, 1636.

said, for instance, that crude ceric sulphate, in the presence of sulphuric acid, is an efficient oxidiser of aromatic hydrocarbons, converting them into aldehydes, quinones, etc., eg toluene into benzaldehyde, anthracene into anthraquinone, etc.

Medicinal Uses.—Cerum is used to a limited extent in therapeuties in the form of cerous nitrate, cerous oxalate, and other organic salts It acts like bismuth as a sedative and is said to be of value in the treatment of sca-sickness, pregnancy-sickness, dyspepsia, and gastric ulcer. The medicinal use of cerium salts is not, however, in very great favour 2

See D.R.P., 158,609 (1905).
 See Eng. Put., 5328 (1884), J. Soc. Chem. Ind., 1889, 8, 301, and Cushny, Text book of Pharmacology and Therapeutics (Churchill, 5th ed., 1910); Brit. Pharmaceutical Codex, 1911.

# CHAPTER XI.

## THE RARE EARTH ELEMENTS (continued).

SEPARATION AND PURIFICATION. HONOGENEITY. DETECTION AND ESTIMATION

Choice of Material. - Monazite in the form of monazite sand is the most abundant of the rare earth nancrals Very large quantities of this material are worked up annually in connection with the gas-mantle industry. For the manufacture of mantles the thornum present in the sand is its most valuable constituent, very little of the cerum and "didymnum" present being required \* Accordingly, even when the residual salts of cerium, lanthanum, etc, are worked up for the production of pyrophoric alloys, etc, large quantities of them are left over, and when it is possible to obtain them in bulk they constitute the most convenient source of the rare earths from the point of view of the scientific investigator. Since, however, monazite sand seldom contains more than one or two per cent of crude yttiia, and frequently contains less than one per cent, it is obvious that very considerable quantities of monazite residues are necessary to obtain a reasonable supply of the earths of the yttirum group, but when they are available they constitute an excellent source of samarium, gadolimum, europium, terbium, dysprosium, and holmium.1 In smaller quantities they constitute the cheapest and most convenient source of ceruin, lanthanum, praseodymnum, and neodymnum Commercial cerum oxalate and "cerum oxalicum medicinale" are essentially monazite residues, and when calcined yield mixtures of oxides containing about lifty per cent of ceria 2

With regard to the other rare earth minerals it may be said that the crude "erbia" of the early chemists is most abundant in venotime, cuvenite, samarskite, and certain Norwegian columbates, while different samples of gadolinite vary considerably in their "cibia" content, that samarskite is comparatively iich in the earths of the terbium group, that the galdbinite earths are rich in yttria and "ytterbia", and that the element celtium, concerning which very little is known, occurs in galdbinite but not in xenotine 5

Extraction of the Rare Earths from Minerals. -The rare earth elements are readily separated from the other elements with which they are associated in nature by taking advantage of the fact that their ovalates are practically insoluble in dilute immeral acids. It is only necessary to obtain the rare earths in solution as the sulphates, intrates, or chlorides in the

James, J. Amer. Chem. Soc., 1913, 35, 235. James and Grant, ibid., 1916, 38, 41.
 Bohm, Zettsch. angere. Chem., 1902, 15, 372.
 J.c. prior to 1878. See the historical section in the preceding chapter (p. 221).

<sup>&</sup>lt;sup>3</sup> Ic. prior to 1878. See the historical section 1
<sup>4</sup> James, J. Amer. Chem. Soc., 1911, 33, 1332.
<sup>5</sup> Urbain, Compt. rend., 1911, 152, 141.

presence of a slight excess of mineral acid, and then, after saturating the solution with hydrogen sulphide, if necessary, and removing the precipitated sulphides, to add a slight excess of oxalic acid. The precipitated oxalates separate at first in an amorphous form and "ball up," so that warm solutions should be used and the first portion of the procipitate vigorously shaken until it assumes the crystalline form before the remainder is precipitated. The amorphous precipitate rapidly becomes crystalline and may then be filtered and washed very readily—On ignition, the oxalates leave a residue of oxides, i.e. the rare earths themselves.

Most rare earth immerals contain thorum, and frequently zirconium is also present. The former is precipitated as oxalate with the rare earth oxalates, but the latter remains in solution, since its oxalate is soluble in oxalic acid and in mineral acids.

The precipitated oxalates are hable to be contaminated with small quantities of iron, aluminium and calcium oxalates, etc., and, when monazite or xenotime has been employed, with small amounts of phosphates. Accordingly, the oxides derived from them by ignition or the hydroxides obtained by boiling them with sodium hydroxide should be dissolved in a slight excess of nitric or hydrochloric acid, and the precipitation with oxalic acid repeated. The alkali and alkaline earth metals may be eliminated by precipitating the rare earths as hydroxides with ammonia, but this is not very convenient on a large scale.

Any remaining impurities usually concentrate in the most soluble fractions, when the rare earths are subsequently separated by fractional crystallisation. Being then concentrated into a small portion of the initial rare earth material, their complete removal offers no difficulty.

The methods that may be employed for opening up rare earth immerals - are as follows.

(i.) Decomposition with Concentrated Sulphuric Acid.—This answers well with the minerals cente, gadolimite, orthite, monazite, and eschyinte. Cente is finely powdered, moistened with water, strifed up into a paste with about two-thirds its weight of concentrated sulphuric acid and heated. The temperature is gradually raised until finally at a dull red heat there is left a residue of anhydrous sulphates. The residue is cooled, pulverised, and slowly added to ree-cold water. The other silicates orthite and gailolimite may be similarly treated, though gadolimite is usually decomposed by heating it with concentrated hydrochloric acid or aqua regia.

Monazite sand requires rather different treatment. It is treated with twice its weight of sulphuric acid, and only heated to 200°-300°. It is necessary to leave a certain minimum amount of sulphuric acid in the residue in order that when it is added to water the phosphates of the rare earth elements may be kept in solution. The mixed phosphates dissolve readily when the residue is stirred slowly into an excess of water, which need not be cooled below the ordinary temperature.

Aschynate 1 is very finely ground and heated with sulphuric acid until excess of the latter has been expelled. The residue is dissolved in ice-cold water and, after twenty-four hours, the liquid is decanted from the residual titanic, columbic, and tantalic acids, etc.

The liquid is treated with excess of ammonia to clummate the considerable quantity of calcium present. The washed precipitate is dissolved in excess

<sup>&</sup>lt;sup>1</sup> Urbain, Ann. Chim. Phys., 1900, [vii.], 19, 206.

of concentrated nitric acid and the solution boiled for several days, more acid being added from time to time to replace that lost by evaporation liquid is then evaporated to a syrip and the residue taken up with water. After several days the clear liquid is decanted from the residue, which contams the last traces of titame, columbic, and tantalic acids, and is extremely difficult to remove by filtration

Ensemble, like aschymite, is a columbo-tantalate. In order that it may be decomposed by sulphune acid it must be very finely ground. According to James, the intric acid process just described for climmating the last traces

of the earth-acids may be omitted

(n) Fusion with Potassium Hydrogen Sulphate —This method was formerly employed to a considerable extent for decomposing rare earth minerals containmg columbium, tantalum, etc., but it is a troublesome process. Samaiskite, aschynite, euxenite, and fergusonite may be treated in this manner. The residue after fusion is dissolved in water and treated as already described in connection with  $\operatorname{reschymite}^{\,2}$ 

(111 ) Fusion with Sodium Hydroride,3 - Various columbates may be decomposed by fusion with twice their weight of sodium hydroxide. The product is cooled, broken up, and allowed to disintegrate in water. It is then acidified with hydrochloric acid, heated, and the columbic acid, etc., removed by

Xenotime (ytterspar) may also be decomposed by this method. The product is boiled with water and filtered. The residue is then dissolved in hydrochloric acid 4

- (iv ) Decomposition with Hydrofluoric Acid 5-This is an excellent, but rather expensive, method for the decomposition of columbates and tautalates, og samarskite, fergusonite, eurenite, æschynite, etc. The mineral is powdered and warmed with aqueous hydrofluoric acid. Columbium, tantalum, titamium, zirconiuif, tingsten, etc., pass into solution and the raio earths are converted into insoluble fluorides. The latter are filtered off and converted into mto insoluble fluorides the neutral sulphates by heating with sulphunic acid, 6 the oxalates are then obtained in the usual manner
- (v) Decomposition by Disulphur Dichloride :- Those rare earth immerals that contain columbic, tantalic, tifamo acids, etc., as then acid components are readily decomposed when heated in a stream of the vapour of disalphur dichloride, S2Cl2 The chlorides of the rare earth metals, etc., remain behind, and the volatile chlorides or oxychlorides of columbium, tantalum, titamium, tungsten, etc., distil over with the excess of disulphur dichloride. The method works excellently with samarskite, fergusonite, aschynite, and currente.

Separation of Thoria from the Rare Earths - It was mentioned in the preceding section that the rare earth oxalates extracted from a immeral

James, J. Amer. Chem. Soc., 1911, 33, 1335.
 See Nilson, Ber., 1880, 13, 1430, Kruss and Nilson, Ber., 1887, 20, 1676, James,

James, J. Amer. Chem. Soc., 1911, 33, 1335.
 See Nilson, Ber., 1880, 13, 1430. Kiuss and Nilson, Ber., 1887, 20, 1676. James, J. Amer. Chem. Soc., 1911, 33, 1336.
 James, J. Amer. Chem. Soc., 1911, 33, 1336.
 A somewhat similar procedure, viz. fusion with sodium carbonate, is recommended by Urbain (Ann. Chim. Phys., 1900, [vii.], 19, 202) for the decomposition of momerite.
 Roscoce, Ber. 1882, 15, 1274. Lawrence Smith, Chem. Acres, 1883, 43, 13, 29, 1885, 51, 289, 304. James, loc. cit., cf. Wolcott Gibbs, Amer. J. Ser., 1864, [ii.], 37, 355.
 Uranium, when present, e.g. in samaiskite, Accompanies the rate earths up to the route.

this point
7 W. B. Hicks. J Amer. Chem. Soc., 1911, 33, 1492, cf. Bailot and Chauvenet, Compt. rend , 1918, 157, 1153, for a similar method, using carbonylchloride, COCl2, as reagent.

frequently contain more or less thorum oxalate.1 It is therefore necessary to discuss the methods by which the thorium may be eliminated.2

(1) The Hydrogen Perorule Method (Method of Wyrouboff and Verneurl).3 -This method is based upon the fact that from a neutral or slightly acid solution of the nitrates of thorium and the raic earth metals, thorium is completely precipitated by hydrogen peroxide in excess. The precipitate is white and voluminous, and has a composition expressed by the formula The O. N.O. all O The precipitation should take place in a solution containing not more than two parts of thora per 100 of solution, the solution should be heated for a few minutes to 85°, the precipitate allowed to settle, filtered, and washed with hot water

When little thorium is present, associated with much cornun, etc., the precipitation of the thorium is complete, but a little cerium peroxide is coprecipitated. On the other hand, when the thorium is in excess, the precipitate is not contaminated with cerum, but a trace of thorium remains in solution

In applying the peroxide method on a large scale, it is advisable, first, to effect an initial concentration of the thorium, next, to remove the thorium by hydrogen perovide, and, finally, to unite the rare earths in the filtrate from the peroxide precipitate with those left over in initially concentrating the thornum. The procedure recommended by Wyrouboff and Verneull is as follows -

The precipitated oxalates t are washed, heated with 10 per cent sodium carbonate to convert them into carbonates, and sodium hydroxide added to precipitate all the thorium that passes into solution. The washed precipitate is dissolved in the minimum of hydrochloric acid, and a suspension of barrium perovide is stirred into the solution a little at a time until a sample of the clear liquid no longer gives a precipitate with pure hydrogen peroxide. The filtered liquid contains the bulk of the rare earths, free from thorium, the precipitate contains all the thorium, contaminated with 20 to 30 per cent of other earths The precipitate is washed, dissolved in cold hydrochloric acid, and the barium removed by a slight excess of sulphune acid. The filtered solution, which should not contain more than 15 per cent, of free acid, is treated with ovalic acid, the precipitated ovalates thoroughly washed, and then treated with a very concentrated solution of ammonium carbonate (to which sufficient ammonia has been added to secure a solution of the normal carbonate) The thornun oxalate passes into solution completely, together with about 7 per cent of its weight of rare earth oxalates The residual oxalates are free from thorium. To obtain the rare earths still associated with the thorium, the cold filtered solution is decomposed by the addition of sodium hydroxide, the precipitate thoroughly washed by decantation, and dissolved in the minimum excess of cold mitric acid. The solution is diluted and the thorium removed by treatment with excess of hydrogen peroxide

See also the section on "Occurrence" in Chap X.

<sup>&</sup>lt;sup>2</sup> No attempt will be made to discuss the purification of thoria; for that the reader is referred to Vol. V.

<sup>3</sup> Wyrouboff and Vernenil, Bull Son chem, 1897, [111], 17, 679, 1898, 19 219, Compt. read., 1898, 126, 340, 127, 412, Ann Chem. Phys., 1905, [viii.], 6, 484, cf. Cleve, Bull Soc. chem., 1885, [11.], 43, 53, Lecoq de Boisbaudtan, Compt. read., 1885, 100, 805

<sup>4</sup> In working up cerum minerals containing a few per cent of thorium, the latter is completely precipitated when sufficient oxalic acid is gradually added to the well-stried acid solution of the sulphates to precipitate one-half of the rare earths present.

All the filtrates containing rare earths devoid of thorium are united, and the rare earths precipitated with oxalic acid in a slightly acid solution

(11) The Ammonium Oxalate Method (Bahr's Method) 1-The mixed oxalates are treated with a hot concentrated solution of animonium oxalate, in which thorium oxalate is readily soluble. The separation of a little thorum oxalate from excess of rare earth oxalate in this manner requires several repetitions of the treatment, and is not a sharp separation, for notable quantities of the rare earth oxalates dissolve, particularly the oxalates of the yttrum group This method has been employed commercially.

(m) The Throsulphate Method (Method of Chydenius) 2 - A dilute, neutral (or slightly acid) solution of the chlorides is boiled with an excess of sodium thiosulphate, when basic thorium thiosulphate is precipitated. This method is frequently employed in the separation of small amounts, ey. in analytical

chemistry (see later, p 371)

(iv.) The Schacic Acid Method !- A slight excess of a hot, concentrated solution of sebacic acid is added to a boiling, neutral or faintly acid solution of the mixed intrates Thornum sebacate is precipitated, filtered, and washed with hot water

Sebacic acid may be prepared from castor oil

(v) The Lead Carbonate Method 1 A dilute and nearly neutral solution of the nitrates is prepared, and an excess of pine, moist, precipitated lead carbonate is stirred into the solution. The solution is allowed to stand for some hours, with occasional shaking. The thorium is completely precipitated, together with zircomum and any cerum present as ceric nitrate

(vi) Other Methods -It has been proposed to precipitate the thorium present by means of sodium azide," or as the fumarate," meta-nitrobenzoate,8 pyrotartrate, 9 iodate, 10 pyrophosphate, 11 hypophosphate, 12 etc 13

1 Bahr, Manalen, 1864, 132, 227, Bunsen, Pong Annalan, 1875, 155, 380, Glaser, J. Anno. Chem. Soc., 1894, 18, 782, Hintz and Weber, Zeitsch. anal. Chem., 1897, 36, 27, 213, 676, Urbain, Ann. Chem. Phys., 1900, [vii ] 19, 181.

2 Urdenius, Pong Annalan, 1863, 119, 46, Hermann, J. pract. Chem., 1864, 93, 106, Drossbath, Zeitsch. angew. Chem., 1901, 14, 655. Hauser and Writh, that, 1909, 22, 484; Hintz and Weber, loc. cit., Glaser, loc. cit., and Chem. Zeit., 1896, 20, 612, Johnstone, J. Soc. Chem. Ind., 1914, 33, 55.

3 T. O. Smith and C. James, J. Amer. Chem. Soc., 1912, 34, 281. See also Knoffer, D.R.P., 266, 459 (1912).

4 Giles, Chem. News., 1905, 92, 1.

5 Hence, if it is not desired that any cerium should be precipitated, care must be taken to ensure the absence of ceric initiate. For this purpose the dilute solution may be saturated with hydrogen sulphide, and the excess of gas removed by boiling, or sulphinous acid may be used.

with hydrogen sulphide, and the excess of gas removed by builing, or sulphinious acid may be used

8 Dennis and Kortright, Amer Chem J, 1894, 16, 79, Dennis, J. Amer Chem. Soc., 1896, 18, 947. Wyroub fland Verneuil, loc et

7 Metzgri, J. Amer Chem. Soc., 1902, 24, 901

8 Neish, ibid., 1904, 26, 780, Kolb and Ahrle, Zeitsch ungew Chem., 1905, 18, 92.

9 Smith and James, J. Amer Chem. Soc., 1912, 34, 281

10 R. J. Moyer and Speter, Chem. Zeit., 1910, 34, 300, R. J. Meyer, Zeitsch anorg Chem., 1911, 71, 65.

11 Carney and Campbell, J. Amer Chem. Soc., 1911, 36, 1134, and D. R. P., 286,087 (1914).

12 Koss, Chem. Zeit., 1912, 36, 686; Rosenheim, ibid., 1912, 36, 821, Witth, ibid., 1913;

37, 773, D. R. P., 268,866

19 Lecoq de Bushandian. Compt. read., 1831, 99, 525. Urbain, Bull. Soc. chem., 1896, 15, 338, 347, Ann. Chim. Phys., 1900, [vii.], 19, 184, Schutzenberger and Boudouard.

15, 338, 347, Ann Chun Phys., 1900, [vii.], 19, 184, Schutzenberger and Boudouard, Compt. read, 1897, 124, 481, Chavastelon, ibid, 1900, 130, 781, Giossmann, Zeitsch. anorg (hrm, 1905, 44, 229; Batčk, ibid., 1905, 45, 87, Muthmann and Baun, Ber., 1900, 332, 2028; Miss Jellerson, J. Amer. Chem. Soc., 1902, 24, 540; Haitwell, ibid., 1903, 25, 1128; Smith and James, ibid., 1912, 34, 281.

Should any trace of thorium be left with the rare earths after one of the preceding methods has been used to effect its removal, it will be subsequently found in the most soluble fractions of the yttrium group if the separation of the earths is commenced by the double sulphate method, using sodium sulphate as precipitant, and the general procedures given later are followed (see p. 357).

SEPARATION OF THE RARE EARTH ELEMENTS FROM ONE ANOTHER. METHODS AVAILABLE. FRACTIONATION CONTROL OF FRACTIONATION.

The preparation of pure certa from a mixture of rare earths is a comparatively simple operation that may be carried out by methods closely resembling the ordinary analytical methods of separation. The preparation of any other rare earth, however, is a tedious process, the ordinary methods of separation being unavailable. In the following account the nature of the separation processes is first outlined, and an indication given of the manner in which they are applied in practice. The actual processes are then enumerated and the application of the more useful of them briefly outlined.

Methods Available. In ordinary analytical chemistry the problem of separating the metals present in a mixture is accomplished by converting each in turn into an insoluble or volatile compound to the exclusion of those accompanying it. Each separation may thus be effected completely, or almost completely, in a single operation. In the case of the raise earth elements, however, the most that can be done in one such operation is to effect a partial separation, i.e. to divide the primitive mixture into two parts containing the raise earths in different proportions. These two parts have then to be further subdivided, and the process of subdivision carried on for a very long time, portions similar in composition being united at appropriate intervals. Such a method of separation is spoken of as a fractionation, and the various parts into which the initial mixture is divided are termed fractions.

The methods of fractionation, which most closely resemble the methods of ordinary analytical chemistry, are as follows:—

(1) Fractional volatilisation.

(ii.) Fractional decomposition of salts by heat.

(iii) Fractional precipitation.

Fractional volatilisation has so far only been employed in effecting the partial separation of ytterbium and lutecium, and need not be further discussed.

The methods of fractional decomposition and precipitation were much employed by the early workers on the rare earths. The former is of little value except in one instance, the decomposition of the intrates of the yttrium group. Both exploit the fact that the rare earths, although all fairly strong bases, exhibit slight differences in their strengths. In the case of fractional precipitation the results obtained also depend upon the relative solubilities of the sparingly soluble compounds that are precipitated, in most cases, however, the solubilities of these compounds do not differ widely among themselves, so that the influence of solubility is of very little account. The weakest base precipitates first and the strongest last. This holds good whether the rare earths are separated by precipitating the hydroxides, basic salts or normal salts; for example, the yttrium compound is the last to precipitate, whether the yttrium group is precipitated as hydroxides, ferre-

cyanides, iodates, stearates, or basic nitrites. A knowledge of the relative strength of the rare earths is therefore of importance for fractional decomposition or precipitation; the relative strengths have been already discussed (p 257).

The method to be pursued in effecting fractional decomposition or precipitation will be evident after reading the description of fractional crystal-lisation, given below, and the short account of the "decomposition of the nitrates," given later (p 353) There is little doubt that the precipitates obtained in fractionating by precipitation are usually solid solutions or mixed crystals.

Fractionation involves the frequent repetition of the same operations. It is accordingly very desirable that the technique of the process should be as simple as possible, in order to economise time, minimise the risk of accidents, and prevent the continual loss of small quantities of material, Methods of fractionation that involve filtrations, ignitions, and techous evaporations should therefore be employed only in special cases, "fractional precipitation to be of value must be very rapid." In general, it is much better to resort to a method of separation which finds no counterpart in ordinary analytical procedure, namely, fractional crystallisation of corresponding isomorphous salts from a suitable solvent. The salts should preferably be very soluble in the hot solvent and separate on cooling in compact crystalline masses. The most convenient solvent is water. The only operations involved consist in dissolving crystals in mother liquors, occasionally concentrating the solutions, and decanting mother liquors from crystals.

Fractional Crystallisation.—The utility of this method for separating the rare earth elements is due to the fact that the corresponding salts of these elements are usually isomorphous and capable of forming mixed crystals. If two salts are not isomorphous, then fractional crystalbaction permits, according to circumstances, of the isolation only of the least soluble or most abundant of them from a mixture. With isomorphous salts, however, it is possible, by a series of skilfully conducted crystalbactions, to separate them in a state of purity, step by step, in the order of their solubilities. The experimental procedure may be varied somewhat to suit a particular case, but in general is as follows:—

The initial mixture of salts (1) is crystallised from the solvent, about half the mixture being left in solution. Crystals (2) and mother liquor (3) are thus obtained and separated. Each in turn is crystallised. The crystals (2) give place to crystals (4) and mother liquor (5); the mother liquor (3) becomes divided into crystals (6) and mother liquor (7). Liquor (5) and crystals (6) are united into one fraction, and this, together with crystals (4) and liquor (7), constitute the three fractions into which the material is now divided. By further crystallisation, crystals (4) yield crystals (8) and liquor (9); fraction (5, 6) gives crystals (10) and liquor (11); and liquor (7) supplies crystals (12) and liquor (13). The following combinations are then effected: (9) with (10), (11) with (12); the method being to pour liquor (13) into a fresh vessel, then pour liquor (11) on to crystals (12), and finally to pour liquor (9) on to crystals (10). Resolution into four fractions is thus accomplished. The subsequent operations, as well as those just described, are sufficiently indicated in the accompanying diagram (fig. 34). In this

<sup>1</sup> James, J. Amer. Chem. Soc., 1908, 30, 184.

diagram even numbers represent crystals and odd numbers denote mother liquors; the arrows indicate how each fraction is subdivided, and the juxtaposition of two numbers represents the combination of two fractions into one.

The method of proceeding with the work of subdivision should now be clear. By this process the initial mixture is resolved into some twenty or thirty fractions, according to the nature of the mixture. The first stage of the fractionation is then finished

During the second stage of the process the number of fractions is maintained constant. Suppose, for instance, that twenty fractions have been obtained. They are numbered from (1) to (20), beginning with the least and ending with the most soluble fraction. Each is crystallised (with readily

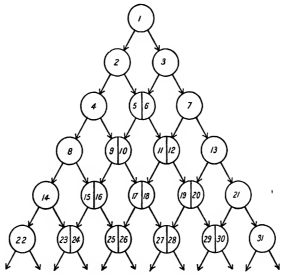


FIG. 31 -To illustrate fractional crystallisation.

soluble substances it is usually best to allow the greater part of the solid to separate out, say, three-fourths of the whole). The mother liquor from (20) is pointed into a new vessel (21), the liquor from (19) poured into (20), that from (18) into (19), and so on until the liquor from (1) is poured into (2). From (2) to (20) the vessels are heated until the crystals dissolve in the liquids, and are then allowed to cool slowly. Fresh solvent is added to (1) for the crystallisation, but (21) is left alone. After crystallisation has occurred, the previous process of decandation is gone through, commencing as before by pouring the liquor from (20) into (21). For the next crystallisation a further quantity of solvent is added to (1), but fraction (21) is still untouched.

After one or two more series of crystallisations, fraction (1) has dwindled to a very small amount. It therefore misses the next crystallisation. After the decantation of mother liquors has been performed, fraction (1) is then

added to the crystals in (2), which thus becomes the new "head" fraction. By this time, however, the accumulation of mother liquors in (21) has produced a new fraction equal in size to the others, and from this time onwards it becomes the new "tail" fraction and takes part in the crystallisations.

By operating in the preceding manner, the fractions (2), (3), (4), etc., disappear in turn, and new fractions (22), (23), (24), etc, are successively added, throughout the entire process, therefore, the number of fractions remains constant. It is generally desuable to maintain the fractions equal in size throughout the series of crystallisations, and, in effecting this, some little skill is required to counteract the tendency of the middle fractions to grow at the expense of those at the two ends. Care should also be taken to exclude dust, etc., and if the crystallisations are carried out in Jena glass flasks, as recommended by Urbain, the flasks must not be more than half filled, otherwise expansion of the crystals not infrequently leads to breakages 1

Serial Order of the Rare Earth Elements. Fractional crystallisation of salts of the rare earth elements usually amounts to the separation of the components of a mixture of isomorphous salts. Consider the case of three such salts, A. B. and C. the solubilities of which increase in the order specified. The prolonged fractional crystallisation of a unixture of these salts leads in general to the following result - The first few "head" fractions consist of pure A, the next fractions in order consist of mixtures of A and B, the percentage of B mereasing as the most soluble fractions are approached then follows a series of fractions consisting of practically pure B; the next series of fractions contain both B and C, the percentage of C increasing in the direction of the more soluble fractions, and, finally, a number of fractions of pure C are obtained 2. In short, the salts are spread out and separated in the order of then solubilities, a number of unxtures or intermediate fractions being interposed between each two consecutive salts

The separation of two rare earth salts by an assigned method requires the presence of a certain number of intermediate fractions. The efficiencies of different methods of separation may be regarded as inversely proportional to the number of intermediate fractions they necessitate. In order to continue with the separation of the salts contained in a series of intermediate fractions, it is necessary to increase their number by the usual method of subdivision and then to continue the fractionation. The intermediate fractions may thus be diminished in quantity, but not in number

In the case discussed it happens that while A has only to be separated from more soluble, and C from less soluble impurities, B has to be separated from a less soluble and also a more soluble impurity In such a case as that of B it is often more difficult to eliminate the less soluble impurity than the more soluble one. The way out of this difficulty, however, is fairly obvious; after (' has been eliminated as completely as possible, the still impine salt B is converted into a new salt B so that the order of mereasing solubility

<sup>1</sup> For further information on fractional crystallisation, see Urbain, J. Chem. phys., 1906, 4, 36. Auer von Welsbach, Monatsh., 1885, 6, 477.

2 This assumes that none of the salts is very scarce in comparison with the others. There is usually no advantage gained by increasing the number of fractions beyond twenty or thirty, and if, say, the salt B constitutes only one-hundredth of the mixture, the state of affairs mentioned in the text obviously cannot be realised. The salt B, however, accumulates is the mixture of the sales see that after fractionation has been unorceded with for in the middle fractions of the series, so that after fractionation has been proceeded with for a sufficient length of time end fractions of pure A and C may be removed. The intermediate fractions may now be increased in number, and finther quantities of A and C removed, until eventually B forms a considerable proportion of the fractions left.

is now C', B', A'. Further fractionation of B' then eliminates A' in the "tail" fractions 1

Similar considerations to the preceding apply when the mixtures to be fractionated contain not merely three, but a considerably greater number of salts In separating complex mixtures, however, the procedure adopted is first to effect a preluminary separation into groups of salts, and then to treat each group apart from the others.2

From a study of the various salts that have been utilised for the purpose of fractional crystallisation it has been found that the order of separation, or serial order, of the rare earth elements is usually as follows -

La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, Dy, Ho, Y, Er, Tm, Yb, Lu (Ct)

Hence, for example, a needymium salt that has been freed by fractional crystallisation from all but the last traces of praseodymium and samarium

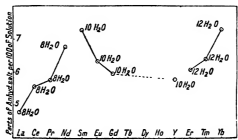


Fig. 35. - Solubilities of the 1:4 2-bromonitrobenzenesulphonates. of the rare earth metals at 25°.

salts, its neighbours in the series, will be even more free from lanthanum, europum, gadolinium, etc.

It is generally assumed that the serial order represents not only the order in which the rare carth salts separate, but also the order of solubility of the salts It is not certain, however, that the separations always do occur in the order of the solubilities of the salts, although this usually appears to be the case.4 It is noteworthy that with the exception of yttrium the serial order of the elements is the order of increasing atomic weights, and thus ranges the terbium group between the cerium and yttrium groups. In the majority of cases the lanthanum salt is the least soluble of the series; occasionally, however, it is the most soluble

The separation of the rare earth salts by crystallisation does not always follow the serial order. It is, perhaps, not to be expected that this order

<sup>&</sup>lt;sup>1</sup> Urbain, Ann. Chim. Phys., 1900, [vii.], 19, 227-29; Baxter and Chapin, J. Amer. Chem. Soc., 1911, 33.-1.

<sup>2</sup> This often involves changing the method of fractionation, and in doing so each fraction should be separately converted into the new salt if the order of separation remains the same; otherwise much of the advantage of the previous seq sation is lost.

<sup>3</sup> Urbain and Lacombe, Chim. News, 19(4, 90, 319, Urbain, J. Chim. phys., 1906, 4, 64; Chem. News, 1909, 100, 73, Seventh Int. Conq. Appl. Chem., 1909, Sect. X., p. 94.

<sup>4</sup> Thus, prascodymium bromate is tather more soluble in water than yttrum bromate, and yet when the mixed bromates of yttrium, prascodymium, crbium, and thulium are fractionated, the prascodymium salt rapidly accumulates in the head fractions in advance of the yttrium (James, J. Amer. Chem. Soc., 1911, 33, 1841). See also the data on p. 269.

would be followed when the salts are not isomorphous and of the same type. The 1:4·2—bromontrobenzenesulphonates¹ are a case in point, as they fall into three groups according to their degrees of hydration. The solubilities have been already given (p. 281); they are plotted against the senial order in fig. 35. It is clear that fractional crystallisation of a complex mixture of these rare earth salts will not effect their separation in serial order. Even when the salts are isomorphous, however, there are cases in which the serial order is not observed. The ethylsulphates (p. 278) afford one such instance, the solubilities diminishing from lanthanum to neodynnum, and thereafter increasing. The bromates (p. 256) and nitrates (p. 267) are analogous to the ethylsulphates, in the case of the bromates the europium salt is the least soluble, and in the case of the initiates, which crystallise from nitic acid as the pentahydrates, the least soluble initiate is the salt of gadolimum.

Separating Elements.—In discussing the atomic weights of the rare earth elements it was pointed out that a number of hismath salts are iso-

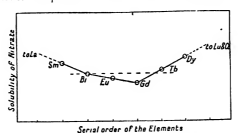


Fig. 36 —Solubilities of the initiates of the rare earth elements.

morphous with the corresponding salts of the rare earth elements (p. 234). If, then, a quantity of the bismuth salt is added to a suitable mixture of rare earth salts, and the whole submitted to fractional crystallisation, the bismuth salt eventually separates out at a definite point in the series, between two of the salts of the rare earth elements. If the crystallisation is continued until a number of fractions of chemically pure bismuth salt have been obtained, a quantitative separation of the rare earths has been achieved. This ingenious method of using bismuth as a separating element is due to Urbain and Lacombe 2

Bismuth occupies a definite position in the serial order, namely, between same rum and europaum, and hence, with mixtures conforming with the law of serial order, it is only possible to produce a quantitative separation at this one point in the rare earth series. When the mixture does not separate in that mainer, however, a different point of separation becomes possible. Only one instance has yet been worked out, namely, the case of the mirates,  $M(NO_3)_3$   $5H_2O$ . When the solubilities of these salts are plotted against the serial order of the elements, the result is screewbat as shown in fig. 36. It will be clear that not only may a quantitative separation be made between samarium and europium if the rare earth elements from lanthanum to

Katz and James, J. Amer. Chem Sec., 1013-35-872.
 Urbain and Lacombe. Compt. rend., 1103-137, 192, 1904-138, 84, 627, 1166; Chem. News, 1903, 88, 295, 1904, 89, 52, 277, Urbain, J. Chem. phys., 1906, 4, 40, 105.

gadolinium only are present, but a similar separation may be made between gadolinium and terbium in mixtures that contain the elements from gadolinium to celtium in the series 1

The application of bismuth as a separating element, when made in counection with the fractionation of the double magnesium nitrates, results in the quantitative separation of samarium from europium, ie the complete climination of intermediate Sm Eu fractions. This is not, however, the only beneficial result that is achieved. The samarium and bismuth salts are miscible to a considerable extent in the solid state, but the neodymium compound is only taken up to a very slight extent, and the praseodymum, cerum, and lanthanum compounds not at all by the bismuth salt. Accordingly, after the bulk of the lanthanum, cerium, praseodymum, and neodymum has been removed in the ordinary way (and this may readily and quickly be effected), the application of the bismuth process leads to the rapid elimination of the remainder of these elements from the samarium, intermediate Nd-Sm fractions being reduced to a very small amount. In a somewhat smular manner the separation of enropium and gadolinium is greatly facilitated, although in this case each salt is readily miscible with the salt of bismuth, four intermediate Eu-Gd fractions only are necessary, instead of eighteen when the bismuth salt is omitted (see p. 319) 2

When the simple nitrates of gadolinium, terbium, dysprosium, etc., are fractionated in the presence of bismuth nitrate, the intermediate Gd-Tb fractions may be reduced practically to zero, and the intermediate Tb-Dy fractions to a small amount

Bismuth may be employed as a separating element in a different manner from the preceding. In fractionating the rare earths, mixtures are sometimes obtained comparatively rich in samarium, and especially in gadolinium, but containing so much of the yttrium group that the samarium and gadolinum cannot be removed from them by crystallisation as the double magnesium nitrates Now the members of the yttrium group do not form double magnesium intrates. If, then, bismuth magnesium intrate is added in sufficient amount to allow the crystallisation to commence, and more added from time to time to the most soluble fraction, the bismuth salt accumulates quite rapidly in the "head" fractions, carrying with it practically the whole of the samarium and gadolinium salts 3

The great value of bismuth for the preceding purposes lies in the fact that when the necessary rare earth separations have been effected by its aid, it may be rapidly and completely eliminated by simple analytical processes Unfortunately, in its isomorphism with the rare earth metals bismuth appears to be unique among the common elements. Occasionally, however, one rare earth element may be used as a separating element for two others

As an example, the separation of lanthanum, prascodymum, and neodymum by the crystallisation of the double ammonium intrates from nitric acid may be discussed. In this process the lanthanum salt separates in the

<sup>&</sup>lt;sup>1</sup> Urbain, Compt rend., 1909, 149, 37.
<sup>2</sup> From the results of Eberhard's investigations of the arc spectra of Urbain and Lacombe's Sm. Eu Gd fractions it may be stated that (i.) the separation of Sm and Eu is quantitative, (ii) pure Eu may be obtained, (iii) Sa may be completely freed from Eu, and (iv) Gd may be obtained quite free from Sm but containing traces of Eu (Eberhard, Zeitsch anny Chem.,

<sup>1905, 45, 374).

3</sup> Urbain and Lacombe. Compt rend., 1904, 138, 84; Chem. News, 1904, 89, 52; Urbain, J. Chim. phys., 1906, 4, 121.

"head" fractions fairly readily, before the praseodymium and neodymium have separated to any considerable extent, and once the bulk of the lanthanum has been thus eliminated, the separation of the other two elements proceeds extremely slowly and it is difficult to induce crystallisation. Further separation of the praseodymium and neodymium may, however, he brought about more rapidly if part of the separated lanthanim is returned to the final mother liquor and again crystallised through the series of fractions. It is better, however, to add cerous ammonium intrate, since the subsequent elimination of cerium is much easier than the removal of the lanthanium? In a somewhat similar manner, James has utilised praseodymium bromate in effecting the separation of the bromates of erbuin and thillium.

In the preceding example, the use of lanthanum or cernin resembles the use of bismuth in facilitating the separation of europium and gadolimum It should be possible, however, to realise the quantitative separation of two rare earth elements by a third in the same way that bisniuth effects the separation of samarium and europium. Thus, suppose a mixture of the salts of three rare earth elements A, B, C be fractionated, the solubilities of the salts increasing in the order A, B, C and the salt of B being in excess of the others, until several consecutive middle fractions consist of practically pure The less soluble fractions will then be A-B mixtures practically free from C, and the more soluble fractions will be B-C mixtures free from  $\Lambda$ Now, by a proper choice of salts the solubilities of which do not follow the serial order from lanthanum to celtum, it may be possible to separate quantitatively each of the binary mixtures A-B and BC by the bismuth magnesium nitrate method, and, if so, the separation of A from C will have been accomplished quantitatively by the aid of B. James and Bissel have obtained good evidence that neodymnum may be used in this manner in effecting a quantitative separation of gadolimium and terbium bromates, in this case A, B, C are gadolimium, needymium, and terbium, respectively, and the A-B and BC mixtures may be quantitatively separated by the bismuth

Control of Fractionation.—It is necessary from time to time to follow the progress of the fractionation. The methods utilised for this purpose are (1) measurement of chemical equivalents, (n) measurement of magnetic susceptibilities, and (iii) observations of various types of spectra.

(i) The "Chemical Equivalent" Method—It is possible to give a perfectly definite meaning to the term "equivalent" when applied to a mixture of rare carths, viz, the number of parts by weight of the mixture that react with one equivalent of an acid, and the experimental determination of the equivalent may be carried out by any of the methods that have been previously described for the determination of the atomic weights of the rare carth elements (p. 239). It is usual, however, to express the result of the measurement, not in terms of an equivalent, but rather as the fictitious "atomic weight" corresponding to it.

Suppose, now, that a rare earth mixture has been resolved into a number

<sup>&</sup>lt;sup>1</sup> Auer von Welsbach, Monath., 1885, 6, 477, Schottlander, Ber., 1892, 25, 378, <sup>5</sup>69.
<sup>2</sup> Von Scheele, Zeitsch annig Chem., 1898, 17, 310, cf. Fert and Pizibylla, Zeitsch, annig Chem., 1905, 43, 202, and Edmarn and Wirth, Innahn, 1908, 361, 190, who use cerous magnesium mitrate to assist in the separation of the double magnesium nitrates of lanthanum, pracedymium, and needymium.

James, J. Amer. Chem. Soc., 1911, 33, 1341.
 James and Bissel, J. Amer. Chem. Soc., 1914, 36, 2060.

of fractions. Frequently, the order of separation of the rare earth elements corresponds with the order of their atomic weights. If, then, the extreme atomic weights of the elements in a mixture are Wa and Wa, the fictitious "atomic weights" of the two end fractions continually approximate to the values Wa and Wa, and the "atomic weights" of the intermediate fractions vary regularly from the one extreme to the other. The manner in which these "atomic weights" vary as fractionation proceeds may be described most simply, perhaps, in terms of a graphical construction, in which the numbers of the fractions are plotted as ordinates and the corresponding "atomic weights" as abscisse, a curve being then drawn through the points. The curve has at first an upward or downward trend throughout its entire length, but as fractionation proceeds, indications of horizontal portions appear in the curve and gradually become more and more pronounced. As a rule, each horizontal region corresponds to a series of fractions of a pure rare earth compound, so that the atomic weights Wa, WB, Wy Wω, corresponding to the horizontal portions of the curve, are the atomic weights of the rare earth elements present in the mixture. Occasionally, however, a horizontal region corresponds to the isolation of a mixture that cannot be resolved by the particular process that is being employed.

Even when the rare earth elements do not separate in the order of their atomic weights the final curve will consist of a number of horizontal regions joined together by intermediate curved portions, its evolution, however, from a simple starting-curve cannot be followed in the preceding manner This is the case, for instance, when the yttrium earths are fractionated by crystallisation, the atomic weight of yttrium being much less than that of any other rare earth element.

In any case, however, the "atomic weight" method of control is used in conjunction with the spectroscopic method. Valuable information may be obtained by it, but, owing to the fact that the atomic weights of any two successive elements in the serial order are very close together,1 the method is not very sensitive.2

(11 ) The Magnetic Susceptibility Method - The rare earths differ very much among themselves in the values of their magnetic susceptibilities. For instance, the values for gadolmia and europia are in the ratio of 5 to 1. Determinations of magnetic susceptibilities therefore provide a much more sensitive means of controlling the course of a fractionation than measurements of chemical equivalents. Moreover, by means of Curic and Cheneveau's magnetic balance it is possible to determine a magnetic susceptibility in a few minutes; the method is therefore extremely practical. Its utility was pointed out by Urbain and Jantsch.3

<sup>1</sup> Yttrum being excluded.

The oraliste method of p 242 is, perhaps, the one that has been most frequently employed in determining the "atomic weights" corresponding to the various fractions. Like practically all the other methods the "weight of oxide" is one term of the latio measured Since, then, cens is a dioxide, it is necessary to chiminate cerium before commencing the fractionation, or else with each "atomic weight" measurement to determine the percentage of cens in the oxide. Similar difficulties are likewise encountered owing to the fact that the oxalates of praccodymium and terbium give peroxides when calcined and not sesquiovides, either the peroxides may be reduced in hydrogen or their peroxide oxygen determined iodimetrically. The "atomic weight" method is accordingly of very little value

so far as the cerum group is concerned

<sup>3</sup> Urbain Compt. rend, 1908, 146, 406, 922; Urbain and Jantsch, thid, 1908, 147, 1286. Urbain, thid., 1908, 149, 37, 1910, 150, 913; 1911, 152, 141, Blumenfeld and Urbain, thid., 1914, 159, 323.

In putting this method into practice the oxalates should be precipitated from the fractions and calcined to oxides just previous to the actual measurements of the susceptibilities. The numerical data for the pure earths are

given on p 257.1

(m) The Spectroscopic Method.—The salts of nine of the rare earth elements give absorption spectra in the visible region, and those of one other give a well-defined ultraviolet spectrum. The frequent observation of the absorption spectra of the fractions therefore affords a rapid and extremely valuable method for following the progress of fractionation. Throughout the early stages of the separation, observations with a good pocket spectroscope are sufficient for the purpose; subsequently, however, more precise measurements are necessary. The comparison of the spectra of different fractions must be effected under conditions which are the same for each fraction, or erroneous conclusions may easily be drawn <sup>2</sup>

When a series of consecutive fractions have identical absorption spectra, it is usually, but not invariably, the case that a pure rare earth salt has been obtained. It is not enough to compare the visible absorption spectra, the ultraviolet regions must also be examined, preferably by the photographic

method.

١.

The separation of the elements in the colourless fractions should be followed by studying the spark spectra, which are not so sensitive as the arc spectra and involve no sensible loss of material in their examination. The study of the spark spectra of the coloured fractions is also extremely valuable. These spectra should not be examined, however, until very considerable progress has been made in the fractionation, since the spark spectra of rare earth mixtures are exceedingly rich in lines. The constancy of absorption and spark spectra throughout several consecutive fractions is the best criterion of the isolation of a pure rare earth compound, but it is not initallible. When this constancy of spectral characteristics is attained, accurate atomic weight measurements should be made, in order to obtain additional confirmation.

Finally, observations of the arc and phosphorescence spectra may be made in order to determine the degree of purity of the rare earth salt that has been isolated

The various spectra of the rare earth elements and their compounds are described in some detail in the preceding chapter (pp. 282-312)

It may generally be assumed that a pure rare earth compound has been isolated in the course of fractionation when a number of consecutive fractions are obtained, identical in all respects, that is to say, when all the fractions correspond to the same atomic weight, yield rare earths identical in their physical properties, particularly their magnetic susceptibilities, and possess identical spectral characteristics, quantitatively as well as qualitatively. When, in addition, the fractionation of such a series of fractions is continued by other methods with negative results, the assumption that a pure rare carth compound has been isolated becomes practically certain. The rare earth thus isolated conforms, in fact, to the experimental definition of a pure substance.

In controlling the course of fractionation one other point should be noted. As has been already pointed out, between each two consecutive series of

<sup>1</sup> On the measurement of magnetic susceptibilities, see Curic, Ann. Chim. Phys., 1895, [vii.], 5, 289; Meslin, ibid., 1906, [vii.], 7, 115. Curic and Chemeveau, J. de physique, 1903, [iv.], 2, 796; Poynting and Thomson, Electricity and Magnetism (Griffin & Co., 1914).

2 See pp. 283-7.

fractions of pure rare earth salts there exists a series of intermediate fractions. It is of importance (i) that these should be expanded into a larger number of smaller fractions, and the process of separation continued until eventually the intermediate portions are insignificant in comparison with the pure salts; and (ii) that confirmation can be obtained that the spectral and other characteristics of these intermediate fractions are in harmony, both qualitatively and quantitatively, with the assumption that they are mixtures of the two pure compounds in varying proportions. Without such confirmation, there is always the possibility that the intermediate fractions contain another earth, rare in comparison with those that have been isolated

## SEPARATION OF CERIUM FROM THE OTHER RARE EARTH ELEMENTS.

The comparative case with which cerium may be separated by chemical means from the other rare earth elements is due to the fact that cerium, unlike the others, forms two definite series of salts, one derived from a strongly basic hydroxide, Ce(OH)<sub>4</sub>, the other from a feebly basic hydroxide, Ce(OH)<sub>4</sub>. The latter hydroxide resembles therium hydroxide, Th(OH)<sub>4</sub>, in strength Accordingly, when cerium is separated from the rare earths by any of the usual procedures, the thorium, unless previously removed, accompanies it

A mixture of rare earths containing not more than 50 per cent of ceria dissolves completely in hydrochloric or intric acid, in the latter case the cerium passes into solution almost entirely as ceric intrate. If, however, a mixture of rare earths contains more than 50 per cent, of ceria, it cannot be completely dissolved in either hydrochloric or intric acid, but it can be completely converted into a mixture of sulphates by heating with excess of concentrated acid. In this manner the ceria is converted into ceric sulphate, insoluble in the excess of sulphiric acid, and if the mixed sulphates are dissolved in cold dilute intric acid and boiled with excess of alkali hydroxide, the precipitated hydroxides contain the cerium, mainly as ceric hydroxide. This precipitate dissolves readily in intric acid, forming ceric intrate, etc.

A rate earth uniture containing more than 50 per cent of ceria can be dissolved by heating it with intite and if hydrogen perovide is added from time to time. The ceria then passes into solution as cerous intrate. The initial mixture may also be dissolved in concentrated hydrochloric acid if ammonium or potassium todide is also added. Todino is liberated and the cerium is transformed into cerous chloride. Ceric chloride is not known.

Natural mixtures of rare earths rich in ceria, e.g those extracted from monazite and cente, do not as a rule contain more than 50 per cent of ceria.

(1) The Basic Nitrate-Sulphate Method.<sup>3</sup>—This method is based upon the fact that ceric intrate and sulphate readily hydrolyse in aqueous

Wyrouboff and Verneuil, loc cit.
 Brauner, Trans Chem. Soc., 1885, 47, 879, Monatsh., 1885, 6, 785, Zettsch. anorg. Chem., 1903, 34, 207. Brauner and Batek, bbd., 1903, 34, 103, Wyrouboff and Verneuil, Bull. Soc. chim., 1897, [iii.], 17, 679; 1898, [iii.], 19, 219, Compt. rend., 1897, 124, 1230; 125, 950; 1899, 128, 1331, Ann. Chim. Phys., 1906, [viii.], 9, 333; Steiba, ibid., 1904, [viii.], 2, 193, R. J. Meyer, Zettsch anorg. Chem., 1903, 37, 378. Cf. the following early memoids: Heimann. J. prakt. Chem., 1843, 30, 184; Mailging, Ann. Chim. Phys., 1849, [iii.], 27, 212, Bunsen, Annacien., 1858, 105, 40, 45, Pogg. Annalen. 1875, 155, 230, 366; Holzmann, J. prakt. Chem., 1858, 75, 321, Jahresber., 1862, p. 136; Czudnowicz, ibid., 1860, 80, 16, W. Gibbs, Amer. J. Sci., 1864, [ii.], 37, 352; Eik, Zeitsch für Chem., 1870, [ii.], 7, 100; Jahresber., 1870, p. 319.

solution, producing precipitates of basic salts that may be readily filtered and washed. The necessary procedure is here described on the assumption that the mixed earths contain less than 50 per cent. of cerm, the details are essentially those of Wyrouboff and Verneuil, as modified slightly by Sterba.

The rare earth elements are precipitated as oxalates, and the latter washed, dried, and calcined to oxides. The mixed oxides are added gradually to four times their weight of cold, concentrated nitric acid, the mixture being warmed towards the end of the operation to 50" or 60°. The dark red solution is allowed to stand for twenty-four hours, and then decanted from any slight insoluble residue. The solution is evaporated on the water bath until it is of the consistency of a thick syrup and solidities completely when cooled. The syrup is dissolved in a quantity of water equal to thirty 1 times the weight of the oxides used, I gram of ammonium sulphate added for every litro of solution, the liquid heated to the boiling-point and boiled for about fifteen minutes. A pale yellow precipitate with a faint greenish tint is thus obtained, it consists of basic ceric intrate and sulphate. The precipitate is filtered off and thoroughly washed with a solution containing 50 grams of ammonium nitrate and 10 grams of ammonium sulphate per litte of water, hot water will serve, but the preceding solution is better. Provided the initial evaporation of the intric acid solution was continued to the right point, the precipitate contains practically all the cermin that was present in the liquid as cene intrate A little ceric nitrate and some corous nitrate, however, still remain in the filtrate

Wyrouboff and Verneuil give two methods for chiminating the remainder of the cerum from the filtrate. (a) The liquid is vigorously stirred and a mixture of aminonia and pure hydrogen peroxide is slowly added. The cerum is thus precipitated as peroxide, accompanied by more or less of the other carths. When the filtered liquid only gives a bluish-white precipitate, the addition of ammonia and peroxide is stopped. The precipitate is washed, dissolved in intric acid, and precipitated by oxalic acid. The oxalates are ignited to oxides, and the procedure already described in the preceding paragraph is followed with the oxides — There is thus obtained, by these two treatments, (a) 99 per cent of the cerum in a nearly pure state, (B) about 90 per cent of the other earths free from cerum, and (y) a small fraction consisting of other earths contaminated with a trace of cerium (b) An excess of sodium acetate is added and the liquid boiled, while ammonium persulphate is added little by little. In this manner all the cerium may be precipitated as basic ceric sulphate, and there is thus obtained three fractions (a) the original precipitate containing 85 to 90 per cent of the cermin in a nearly pure state,  $(\beta)$  about 99 per cent of the other earths quite free from cerum; and (y) about 10 to 15 per cent of the cerum (from the persulphate precipitate) containing a few per cent of foreign earths. The fraction ( $\gamma$ ) may be dissolved in nitric acid, evaporated to a syrup, diluted with water, and boiled, when the cernum it contains is precipitated almost completely and practically free from other substances

The great advantages of the basic intrate-sulphate method are that in one operation it separates about three-fourths of the total cerima in an almost pure state, and the precipitate may be purified in the same manner as that by which it was produced (p 337).

<sup>&</sup>lt;sup>1</sup> When operating on large quantities of material, this may be reduced to ten.

When the initial mixture is such that the cerium is most conveniently brought into solution as a cerous salt, a mixture of the nitrates is prepared. The following procedures are then available :-

- (a) The concentrated solution of the nitrates, slightly acidified with nitric acid, is electrolysed in a large platinum dish which serves as the anode. The current density should be 0.5-0.7 ampères per sq. dcm. of anode surface. About 70-80 per cent. of the cerum may be thus converted into ceric nitrate. The solution is treated with dilute ammonia until a permanent turbidity is first produced, when the procedure already outlined may be followed 1
- (b) The nitrates are converted into double ammonium nitrates and the latter dissolved in water (2 litres per 100 grams of double nitrates). The solution is boiled, stirred with steam, and a solution of magnesium acctate and hydrogen peroxide added drop by drop (50 grams magnesium acetate dissolved in 500 c.c. of 2.5 per cent. peroxide per 100 grams of double nitrates), until a filtered sample no longer gives the orange-yellow cerum precipitate. The precipitate contains all the cerium (as basic percenc acetate), but is contaminated with about 4 per cent. of other earths. It is filtered off and washed with boiling water.2

The precipitate is dried at 120° in order to convert it into basic ceric acetate. It is then dissolved in excess of concentrated nitric acid and the solution evaporated on the steam bath till the free acid is almost entirely expelled and the residue has the consistency of a very thick syrup. The cerium is then precipitated as basic ceric nitrate-sulphate by the method already described

There are thus obtained three fractions: (a) nearly pure basic ceric nitrate-sulphate, containing the greater part of the cerum,  $(\beta)$  the filtrate from (a) containing part of the cerum and a little lanthanum, etc; and  $(\gamma)$ the filtrate from the basic percein acetate, free from ceimin, and containing the greater part of the lanthanum, etc.

(c) Wyrouboff and Verneurl's procedure (b) above, using aminomium persulphate and sodium acetate, may be employed Witt and Thiel 8 replace the sodium acetate by precipitated calcium carbonate.

(ii.) The Bromate Method -Potassium bromate may be used to oxidise cerous to ceric salts, and, provided that the solution is kept almost neutral, the ceric salt hydrolyses and basic ceric salt is precipitated.

As the starting material, a nearly neutral, concentrated solution of the nitrates is required. The solution is heated to boiling, and potassium bromate added. As soon as red fumes of bromme make their appearance, two or three lumps of marble are added and the liquid boiled for one or two hours. As soon as the peroxide test (p. 367) on a sample of the filtered liquid indicates that very little cerium is left in solution, the boiling is discontinued, the marble removed, and the precipitate allowed to settle. The liquid is syphoned off and the precipitate washed with 5 per cent ammonium nitrate solution The precipitate thus obtained from a concentrated solution is dense, settles readily, washes easily, and contains very little impurity. It consists

Sterba, loc. cvl
 R. J. Meyer and Koss, Ber., 1902, 35, 672; cf. Popp, Annales, 1864, 131, 859; Erk,
 Zeitsch. für Chem., 1870, [ii.], 7, 100. According to Wyroubolf and Verneurl, magnesium gestate offers no advantages over sodium acetate.
 Witt and Thiel, Ber., 1900, 33, 1815.
 James and Pratt, J. Amer. Chem. Soc., 1911, 33, 1826.

mainly of basic ceric nitrate when only a slight excess of bromate is employed, and mainly of basic ceric bromate when excess of bromate is used.

The filtrate is boiled with a little more potassium bromate and marble until the liquid is quite free from cerum. The precipitate thus obtained contains a little of the other rate earths, and is dissolved in nitric acid and added to the next lot of starting material. The filtrate contains the other rare earths, free from cerium.

(iii) The Permanganate Method 1—When potassium permanganate is added to a hot, neutral solution of cerous intrate, it is immediately bleached, and a brown precipitate appears, consisting of a mixture of ceric hydroxide and hydrated manganese dioxide. After a certain quantity of permanganate has been added, the colour is no longer discharged. The solution is then found to give an acid reaction with litmus and to contain cerium in solution. The addition of a little alkah leads to the bleaching of the permanganate and the precipitation of more cerium, and if the liquid be kept neutral by the addition of alkah, the permanganate colour only ceases to be bleached when all the cerium is precipitated. In putting this method for the precipitation of cerium into practice, zinc oxide, magnesia, calcium carbonate, and sodium carbonate have all been used as neutralising agents. Of these, the last is probably the best to employ—

 $\begin{aligned} 3\text{Ce}(\text{NO}_3)_3 + \text{K} & \text{MnO}_1 + 4\text{Na}_2\text{CO}_3 + 8\text{Ha}_2\text{O} \\ &= 3\text{Ce}(\text{OH})_4 + \text{MnO}_2 \text{2H}_2\text{O} + 8\text{NaNO}_8 + \text{KNO}_8 + 4\text{CO}_2. \end{aligned}$ 

In order to use this method for the separation of ceruum from the rare earths, a neutral solution of the rare earth intrates should be prepared, containing the cerium as cerous intrate. The solution is heated to boiling, and potassium permanganate solution added in small quantities at a time until the red colour just begins to be permanent. At this point the precipitant is changed for a solution of permanganate to which has been added sodium carbonate in the exact ratio KMnO<sub>4</sub>:4Na<sub>2</sub>CO<sub>8</sub>. This solution is added very slowly to the nearly boiling intrate solution. A faint colour is added very slowly to the nearly boiling intrate solution of permanganate should be maintained all the time, a little pure permanganate being added if at any time the colour is bloached When the cerium ganate being added to at any time the colour disappears more slowly after each addition of precentant and the effectivescence is less noticeable. The acidity addition of precipitant, and the effervescence is less noticeable of the solution is their tested from time to time with litmus paper, and small amounts of the permanganate-carbonate mixture or pure carbonate are added until the solution is nearly neutral to litmus and still pink with permanganate. The whole is heated, stirred for ten minutes and filtered hot, the precipitate being washed thoroughly with boiling water. The precipitate is practically free from other care earths, but it should be dissolved in hot concentrated hydrochloric acid, diluted and precipitated with oxalic acid, the cerium oxalate converted into cerous intrate, and the entire process repeated. cerium is then free from other rare earths, but must be separated from manganese.2

News, 1911, 103, 303.

The procedure here described is due to Roberts (loc. cit )

Winkler, J. prakt. Chem., 1865, 95, 410; Stolba, Jahresber., 1878, p. 1059.
 Drossbach, Ber., 1896, 29, 2152.
 Dr. R.P., 113,106 (1903).
 Muthmann and Rolig, Ber., 1898, 31, 1718;
 Muthmann and Weiss, Annalen, 1904.
 331, 1. Bohm, Zeitsch anger Chem., 1903, 16,
 1129.
 R. J. Meyer and Schweitzer,
 Zeitsch anorg Chem., 1907, 54, 101.
 Exposito, Proc. Chem. Soc., 1907, 23, 64; James,
 J. Amer. Chem. Soc., 1908, 30, 982.
 Roberts, Amer. J. Sci., 1911, [iv.], 31, 350.
 Chem. Nucs., 1911, 103, 303.

in the received from precipitate communical acrees of continue in sen made quite neutral. This trace may be removed, together with see of the other earths, by heating with a little permanganate carbons chittion containing more than four molecular proportions of carbonate p noiccule of permanganate.

(iv.) Mosander's Chlorine Method.—In this process the mixe plorides are precipitated as hydroxides, the latter suspended in excess of pidium hydroxide, and a stream of chlorine passed into the cold suspension antil the liquid is saturated. Cerium is left as insoluble ceric hydroxide; the other rare earths pass into solution as chlorides, the solution of the lanthane being effected much more rapidly than that of the "didymia." epitate is thoroughly washed, dissolved in hydrochloric soid, and the process espeated. Six or seven repetitions of the alkali and chlorine treatment are required, so that the process is very tedious, and, unless the liquid is boiled or some minutes at the end of each chlorine treatment, a little cerium is found in the solution. Instead of chlorine, excess of bromine may be added and the whole heated until the excess has been removed (Browning and

Owing to the difficulty of removing the last traces of impurities from the rium by this method, it is advisable to use it for the isolation of nearly pure ric hydroxide and continue the purification by other methods, e.g. (i.) or (v.);
(v.) The Ceric Ammonium Nitrate Method. —Ceric nitrate forms Ith ammonium nitrate an orange-yellow beautifully crystalline double salt, (NO<sub>s</sub>)<sub>4</sub>.2(NH<sub>4</sub>)NO<sub>s</sub>, which is only sparingly soluble in nitric acid. It is of different type from, and is not isomorphous with, the double ammonium: sitrates of the other rare earth elements. Owing to the nature of this salt, is easy to remove the bulk of the cerium from a mixture of crude cerium erths as ceric ammonium nitrate, and to continue the purification of the frium salt thus obtained. The following procedure is perhaps the most selvenient for commencing to work up large quantities of crude cerium in the laboratory, since it allows the greater part of the cerium to be sparated rapidly in a small volume of solution.

three times their weight of concentrated nitric acid. Assuming that ceries estitutes one-half of the oxides, the requisite amount of ammonium trate to form ceric ammonium nitrate is added. The finely-divided amonium salt may be stirred into a hot solution of the nitrates or added inform of a concentrated aqueous solution. On cooling, the bulk of the

1870 (ii.) 7, 100.

jee Henjer and Wirth, loc. ett.; Browning, Cempt. rend., 1914, 158, 1579;
1916, 170, 49. The fact that the hydroxide of lanthanum passes into solution
i finali the hydroxide of "didymium" is of some practical interest.

1916, 1916, 1917, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918, 1918,

cation of the and washed with opposite the first soid.

In fainthe own ammonium nitrate may contain some five per objective. It is divided into several batches; the first is recrystallised.

nitrio soid (S of soid to 2 of water), the second recrystallised from the first, and so on. Several crops of nearly pure amponium nitrate are thus obtained.

(vi.) Miscellaneous.—Several other methods have been prop eparating cerium from the other rare earth elements, e.g. precipitati offic valerianate, fractional solution of the oxides in dilute nitric activating aqueous ammonium chloride, fusion of the nitrates with exceptionassium nitrates at 330-350°, fractional decomposition of the chromat solubility of ceric oxalate in ammonium oxalate, etc.

The Preparation of Pure Ceria.—(i.) From Basic Ceric Nit he solution evaporated on the steam-bath to the consistency of a very ayrup, cooled, dissolved in cold water, and the liquid boiled for 10-15 min that is to say, the precipitate is purified by a process identical with the which it was produced (p. 333), the addition of ammonium sulphate, how being unnecessary. According to Sterba, four precipitations as basic after diphate are sufficient to eliminate the last traces of other rare earths deria, starting with the crude monasite or cerite earths from which thore

The preceding method also serves for the elimination of the impurity from basic ceric sulphate obtained by the persulphate method. It is conserver, answer for the purification of ceria obtained by the permangation method (p. 385), for which purpose the ceric hydroxide is dissolved out of precipitate by strong nitric acid, which leaves the hydrated manging of findissolved. In the first precipitation of the basic ceric all sulphate, however, the requisite ammonium sulphate must be added.

With careful working, the preceding method of purification gives

excellent yield of pure product.

(ii.) From Ceric Ammonium Nitrate.—The salt is recrystallised strice acid four or five times. The method already outlined is a gonomical one for the purpose.

The combined filtrates may be treated in various ways. A convenient methol affortate them on the steam bath to a thick syrup and precipitate the remaining can begin deric nitrate sulphate (p. 332). Little cerium then remains in the filtrate; porate them on the steam bath to a thick syrup and precipitate in the filtrate; basic derio nitrate sulphate (p. 332). Little cerium then remains in the filtrate; minoved by the permanganate method (p. 335), of by any other convenient procipitate. Sulphan Soilach anges. Chem., 1903, 15, 372).

Buomsparte, Compt. rand., 1843, 16, 1008.

Mosander, Phil. Mag., 1843, [iii.], 23, 241; Scherer, Pogg. Annalen, 1842, 5 Minos, And. Chem. Phys., 1849, [iii.], 27, 200; Zachiesche, J. prakt. Chem. 185; Frericha and Smith, Annales, 1878, 197, 331; cf. Gibbs, Amer. J. Sch. 1908, 23, 21

<sup>35.</sup> Freriche and Smith, Anades, 1875, 197, 481; 97, 41000, americal St.

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Watts, C. J. Chem. Soc. 1849, a, 140; Espainte, Proc. Chem. Soc., 1906; 2 Debrity Wishbod: Debray; Compt. Frank., 1883, 96, 828, Chem. None, 18 in fact theres., Amer. Chem. Soc., 1864, 18, 949; Schukkenburger; Chem. Soc., 1864, 1867, 644; 1867, 124, 481; 1894; task, 772, 109d; Wyronbod and Fernand, lot. oil.; Espainto, Proc. ha 50; of Chadricowine, Proc. Mar. 1897, 30, 18; Gibbs, Ion. oil. Anades, 1897, 1898; Mar. Schuke, Chem. None; 1867, 1898; Espointo, 160; oil. Anades, 1867, 1898; Mar. Schuke, 1867, 1899; Ma

This method is also an excellent one for preparing pure ceria, and may obviously be applied to the purification of the precipitates obtained by Mosander's and the permanganate method.

(iii.) From Basic Ceric Nitrate-bromate - The precipitate is dissolved in concentrated mirror acid by the aid of a very little alcohol, the solution diluted with water and boiled after the introduction of some lump marble. The precipitate is washed with ammonium nitrate, converted into anhydrous cerous sulphate, and the latter dissolved in ice-cold water Hydrated cerous sulphate is then crystallised out by warming the solution to 30-40°. Traces of other rare earth sulphates may readily be eliminated in this manner, since cerous sulphate is not isomorphous with them.

Traces of thorum may be eliminated by the recrystallisation of cerous sulphate, but where a solution of ceric nitrate or ceric ammonium nitrate is available, it is preferable to reduce with hydrogen perovide to the cerous state and eliminate the thornum with hydrogen peroxide from a hot, neutral (or feebly acid) solution (p 320).

A cerous salt in concentrated aqueous solution should show no trace of the absorption spectrum of "didymium" when examined through a very thick layer (10-20 cms), and the residual oxide obtained by the ignition of the oxalate, intrate, double intrate, or sulphate (at a white heat), should have a pale yellow colour when cold, entirely free from any suggestion of a salmon, red or brown tint.1

## PRELIMINARY SEPARATION OF THE RARE EARTH ELEMENTS INTO GROUPS.

In commencing the fractionation of the rare earths obtained from a mineral, the procedure employed depends upon the relative amounts of the rare earths present Two cases present themselves—(1.) the cerum group is present to a much greater extent than the yttrum group, and (11) whe versa In the first case the predominating element is cerum, in the second it is yttrum The terbium group never seems to predominate.

Potassium sulphate is the classic reagent for effecting a separation of the cerium and yttrum groups.2 The double sulphates of potassium and the rare earth elements may be classified according to their solubility in a cold saturated solution of potassium sulphate as follows:-

(a) Those practically insoluble (La, Ce, Pr, Nd). (b) Those sparingly soluble (Sm, Eu, Gd, Tb)

(c) Those more readily soluble in that solution than in pure water (Dy, Y, Er, Tm, Yb, Lu, Ct).

It might therefore be anticipated that a sharp separation of the cerum and yttrium groups could be effected by shaking a cold solution of the sulphates with potassium sulphate in excess, and that the bulk of the terbium group would accompany the cerum group. Such, however, is not the case. The precipitated double sulphates are found to be contaminated with a certain quantity of the yttrium group. Further, the filtrate is found

<sup>&</sup>lt;sup>1</sup> On the preparation of pure ceria, see especially Steiba, Ann Chim. Phis., 1901, [viii.], 2, 193, Wyrouboff and Veineuil, ibid., 1906 [viii.], 9, 333; Diossbach, Bei., 1900, 33, 3506, R. J. Meyer, Zeitsch. anorg. Chem. 1903, 37, 378; Brauner and Batek, ibid., 1903, 34, 103, Brauner, ibid., 1903, 34, 707, Noish, J. Amer. Chem. Soc., 1909, 31, 517.

<sup>8</sup> Klaproth, Ann. Chim., 1803, [1.], 49, 256; Berzelius and Hisinger, ibid., 1803, [1.], 50, 245, Berzelius and Gahn, Schweiger's J., 1814, 16, 250, 404; Beilin, Pogg. Annales, 1834, 43, 105; Mosander, Phil. Mag., 1843, [iii.], 25, 241.

to retain a considerable portion of the terbium and appreciable quantities of the cerum group, particularly neodyminm 1 and samarium salts, for the double sulphates of the latter elements are appreciably soluble in the presence of much yttrium sulphate, and, in addition to the solubilities of the double sulphates of the terbium group being augmented, these double salts are only slowly formed.

The double sulphate separation, then, can only be regarded as a process It may conveniently be carried out by adding finely-divided of fractionation potassium sulphate continuously to a cold dilute solution of the nitrates, chlorides, or sulphates contained in a tall vessel and vigorously stirred by a water-turbine, 2 or a hot saturated solution of potassium sulphate may be stirred into a cold solution of the rare earth salts? The whole is allowed to stand, with occasional shaking, until the presence of "didyminin" in the liquid (as indicated by its absorption-spectrum) can searcely be detected. The precipitate is then filtered off and washed with a cold saturated solution of potassium sulphate. The earths in the filtrate are then precipitated either as hydroxides or oxalates.

Instead of using potassium sulphate, the more soluble sodium sulphate has been used, and, indeed, is said to be preferable 4. The finely-powdered solid is stirred into a ten per cent solution of the rare earth chlorides, intrates, or sulphates Care must be taken, however, not to add an unduly large excess of sodium sulphate owing to the possibility of thereby precipitating yttrium as the double sulphate

The preceding method is not very satisfactory when the ceruin group predominates very largely over the yttrium group, eg with the monazite earths, and one of the following alternative processes may be adopted: (a) The potassium sulphate method is used, but the precipitant is added slowly, and in quantity insufficient to precipitate all the cerium group leaving a considerable amount of the needymum in the solution, practically none of the metals of the terbium and yttrium groups are precipitated earths in the filtrate are precipitated as oxalates, converted into double magnesium nitrates, and fractionated from dilute nitrie acid until the final mother liquor refuses to crystallise. The crystals contain the cerium group up to neodymium, the mother liquor contains samarium and the terbium and yttrium groups. The further treatment of this liquor may be carried out according to the procedure described later for working up the final fractions ( $\delta$ ) of the double magnesium untrates (see p. 349)  $^{\circ}$  (b) The entire material may be fractionated by the double magnesium intrate method, the terbum and yttimm groups rapidly concentrating in the most soluble fractions (c) A solution of the sulphates is prepared, a considerable excess of sulphuric acid added, and the liquid warmed slightly. The bulk of the cerium group is thus precipitated as snlphates. The filtrate is partly

Boudonard, Compt. rend., 1898, 126, 900, Bull. Soc. chim., 1898, [iii], 19, 382;
 Ulbain, Bull. Soc. chim., 1898, [iii.], 19, 381; Ann. Chim. Phys., 1400, [vii.], 19, 251.
 Pennis and Chamot, J. Amer. Chem. Soc., 1897, 19, 799
 For modifications of the piecess, see Bettendorf, Annalen, 1891, 263, 164; Maiignec, Ann. Chim. Phys., 1880, [v.], 20, 535. Muthmann and Rolig, Ber., 1898, 31, 1718; Lecoq de Boisbaudran, Compt. rend., 1886, 102, 902, 1003
 Gibbs, Amer. J. St. 1864, [ii.], 37, 354; Delafontaine, Compt. rend., 1881, 93, 63; Smith, Amer. Chem. J. 1883, 5, 44, 73. Drossbach, Ber., 1896, 29, 2472, 1900, 33, 3506; James, J. Amer. Chem. Soc., 1908, 30, 979, 1919, 34, 757.
 See p. 433 and fig. 42.
 James, J. Amer. Chem. Soc., 1918, 35, 235.

<sup>&</sup>lt;sup>6</sup> James, J. Amer. Chem. Soc., 1913, 35, 235.

neutralised with sodium hydroxide. The remainder of the cerium and part of the terbium group are thereby separated as double sulphates. (d) The double carbonate method given later (p. 342) is applied in order to precipitate the bulk of the cerium group, the yttrium and terbium groups remaining in solution 2

When the yttrium group largely predominates in a mixture of rare earths, as, for example, in the earths extracted from xenotime or gadolimite, Urbain strongly recommends the fractionation of the ethylsulphates as being the best method for commencing the separation. The ethylsulphates should be methodically separated into twenty fractions and then submitted to sixty series of crystallisations. The separation does not occur in serial order. The neodymnum salt separates first. The lanthanum salt, the most soluble in the cerum group, is approximately equal in solubility to the gadolinium salt. From gadolimum onwards the serial order holds good. Accordingly, after the fractionation has been carried to the degree just mentioned, the head fractions are examined for lanthanum,4 and all fractions containing that element are removed. They contain the cerium group, europium, gadolinium, and terbium, and their subsequent fractionation is commenced by the double magnesium nitrate method The remaining fractions consist essentially of yttrum ethylsulphate, the least soluble containing in addition terbium, dysprosium, and holmium, and the most soluble erbium, thulium, ytterbium, and lutecium

According to Urbain, the preceding method of fractionation also constitutes the best as yet available for effecting the separation of the terbium and yttrium groups. For this purpose the cthylsulphate fractions remaining after the removal of those containing lanthanum are submitted to further series of fractionations. The more soluble fractions may be set aside as soon as the rare earths extracted from them exhibit a pure white or pale pink tint entirely free from any trace of an orange tint In this manner the yttrium group may be completely freed from the terbum group.5 An excellent alternative is the fractional crystallisation of the bromates (see pp 348, 351, 355-6).

There are two methods, resembling one another closely, whereby a reasonably sharp separation of lanthanum, cerum, prascodymium, and neodymium from samarium, europium, gadolinium, etc., can be quickly effected. The first consists in fractionally crystallising the double magnesium nitrates of the type 2M(NO<sub>3</sub>)<sub>2</sub>.3Mg(NO<sub>3</sub>)<sub>2</sub>.24H<sub>2</sub>O from nitric acid of density 1.3, and

<sup>1</sup> Drossbach, Ber., 1896, 29, 2452; Chem. News, 1896, 74, 274
2 Drossbach, Ber., 1900, 33, 3506.
3 Urbain, Compl. rend., 1898, 126, 835; Bull. Soc chim., 1898, [in.], 19, 376, Ann. Chim. Phys., 1900, [vii.], 19, 184; 1909, [viii.], 18, 274, J. Chim. phys., 1906, 4, 55
4 By means of their spark spectra.
5 The preparation of the ethy sulphates is described elsewhere (p. 278). It is extremely important that the solutions should be free from and. In fractionating, the solutions should never be heated any more than is necessary, since the salts are hable to hydrolyse, to a considerable extent—an occurrence which may be recognised by solutions originally neutral sheeming and. The small amounts of sulphates that may thus be produced in solution are easily reconverted into ethyl-sulphates by double decomposition with barium ethylsulphate. The crystals formed in one fraction are dissolved in the mother liquor from the preceding fraction by vigorous shaking, and heat is only applied when very little solid remains andissolved. The crystals deposited in any operation of crystallisation should occupy about one-third the total volumes of crystals plus liquid. The final mother liquois about one-third the total volumes of crystals plus liquid. The final mother liquois about one-third the total volumes of crystals plus liquid. The final mother liquois about one-third the total volumes of crystals plus liquid. The final mother liquois about one-third the total volumes of crystals plus liquid. The final mother liquois about one-third two the total volumes of crystals plus liquid. The final mother liquois about one-third the total volumes of crystals plus liquid. The final mother liquois about one-third two theorems are assily reconversed to the entire the total volumes of crystals plus liquid. The final mother liquids about one-third the total volumes of crystals plus liquid. The final mother liquids about one-third the total volumes of crystals plus liquid. The final mother liquids about one-third the total volumes of crystals pl

is known as Demargay's Method, 1 the second, due to Lacombe, consists in fractionally crystallising the double manganese nitrates in a similar manner.2 These double salts separate in serial order, and it will be seen! from an inspection of the table of solubilities (p. 269) that in passing from neodymium to samarium there is a large increase in solubility. By either of the preceding methods samarium, enropium, etc., rapidly concentrate in the most soluble fractions, which acquire an orange colour and show the principal absorption bands of samarum

The methods of Demarcay and Lacombe may be regarded as methods for separating the cerum and terbium groups, although samarium, which is always classed with cerum, accompanies the terbium group of elements; they have

other uses, however, as will be seen later '

# THE SEPARATION OF LANTHANUM, PRASEODYMIUM, AND NEODYMIUM.

In order to accomplish this separation, recourse must be had to methods of fractional crystallisation. The separation of praceodymia from needymia by methods depending on differences of basicity seems to be almost impossible to effect.

The method by which lanthanum and "didymium" were first separated was a crystallisation process, utilising the sulphates for the purpose, but the carly workers never carried out the separation in a sufficiently prolonged and systematic manner to resolve the "didymnum" into its components. Subsequently, the method gave place to a basic one, namely, fractional precipitation with ammonia, and thus it happened that the separation of samarinin from "didyminin" preceded by some six years the discovery of prascodymium and neodymium

A Separation by Fractional Crystallisation.—The following salts have been utilised for the purpose .-

(1) Nitrates, 4 M(NO<sub>3</sub>), 5H<sub>2</sub>O.

(ii ) Double ammonium nitrates, M(NO.), 2(NH4)NO3.4H2O.

- (ii) Double sodium intrates, M(NO<sub>4</sub>)<sub>2</sub>, 2NaNO<sub>3</sub> × H<sub>2</sub>O. (iv) Double magnesium intrates, <sup>7</sup> 2M(NO<sub>3</sub>), 3Mig(NO<sub>4</sub>)<sub>2</sub>,24H<sub>2</sub>O. (v) Double manganese intrates, <sup>5</sup> 2M(NO<sub>3</sub>)<sub>3</sub> 3Mig(NO<sub>4</sub>)<sub>2</sub> 24H<sub>2</sub>O.
- (vi) Chlorides, MCl, 6 or 7H, O.

1 Demarcay, Compt. rend., 1900, 130, 1019, 1185.
2 Lacombe, Bull Suc chim, 1901, [iii], 3x, 570, Chem. News, 1904, 89, 277
3 For the utility of the dimethylphosphates see J. C. Morgan and C. James, J. Amer., Chem. Suc., 1914, 36, 10
5 Fractional precipitation of the formates and lactates in actione solution has been siggested by Baim bey, ibid., 1912, 34, 1174.
4 Demarcay, Compt. rend., 1890, 122, 728, 1900, 130, 1019, Fert and Pizibylla, Zeitsch. amorg. Chem., 1905, 43, 202, Baxter and Chapin, J. Amer. Chem. Soc., 1911;

33. 1

8 Mendeléeff, J. Russ. Chem. Soc., 1873., Annalm., 1873., 168, 45., Auer von Welsbach,
Monatsh., 1885., 6, 477. Setzungsber. K. Akad. B. iss. B. ien., 19.3., 112, 11., A, 1037.; von.
Scheele, Ber., 1899., 32., 409., Dennis and Chamot. J. Amer. Chem. Soc., 1897., 19, 799;
Baxter and Chappin, ibid., 1911., 33, 1., Baxter and Stewart, ibid., 1915., 37, 516., Schoots-

Barter and many parties of the state of the 1905, 43, 202.

Lacombe, Bull. Soc. chim., 1904, [111], 31, 570; Chem. News, 1904, 89, 277.
 Baskei ville and Stevenson, J. Amer. Chem. Soc., 1904, 26, 54.

 $\begin{array}{lll} \mbox{(vii.)} & \mbox{Sulphates,}^1 \ M_2(SO_4)_8 \ rH_2O. \\ \mbox{(viii.)} & \mbox{Ethylsulphates,}^2 \ M(C_9H_5 \ SO_4)_3 \ 9H_2O. \\ \mbox{(iv.)} & \mbox{Biomates,}^3 \ M(Bro_3)_8 \ 9H_2O. \\ \mbox{(v.)} & \mbox{Ovalates,}^4 \ M_2(C_2O_1)_1 \ rH_2O. \\ \end{array}$ 

(x1) Metantrobenzeuesulphonates,  $^{5}$  M[C<sub>4</sub>H<sub>4</sub>(NO<sub>2</sub>)SO<sub>3</sub>]<sub>3</sub>· $^{6}$ H<sub>2</sub>O. (x11) Double potassum carbonates,  $^{6}$  K<sub>2</sub>CO<sub>3</sub>·M<sub>2</sub>(CO<sub>3</sub>)<sub>3</sub>·12H<sub>2</sub>O.

(xini.) Picrates, M[C6H2(NO2)3O]3.

The double nitrates, sulphates, ethylsulphates, bromates, piciates, and metantrobenzenesulphonates may be crystallised from water, but the double ammonium, sodium, and manganese nitrates are best crystallised from strong nitric acid, in which they are much less soluble than in water The chlorides are crystallised from concentrated hydrochloric acid, the oxalates from dilute nitric acid, and the double carbonates from aqueous potassium

The sulphates must be dehydrated, dissolved in ice-cold water, and crystallised by warming the solution to 30°-40°. The double carbonate method is carried out as follows -A 20 per cent solution of the chlorides is poured drop by drop into excess of 50 per cent. potassium carbonate, whereby a solution of the double carbonates in potassium carbonate is obtained.5 The double salts are fractionally precipitated by successive additions of water; they separate slowly, and readily form supersaturated solutions The ovalates may be suspended in water, the latter heated to boiling, and nitric acid added with vigorous shaking until complete solution is effected. The liquid is then rapidly cooled, being agitated throughout the process. Another method is to addify strongly a solution of the nitrates with nitric acid, and drop ovalic acid solution into the nearly boiling nitrate solution until the precipitate that first forms no longer redissolves. The liquid is then cooled as before.

The order of separation is as follows, the least soluble being placed first -

Double nitrates La, (Ce), Pr, Nd; Metanitrobenzene-

La, (Ce), Pr, Nd "Di," La La, Nd, Pr sulphonates Ethylsulphates Nd, Pr, La; Ovalates Nd, Pr, La, Bromates Sulphates Nd, Pr, (Ce), La; Double carbonates La, Pr, (Ce), Nd. Nitrates

<sup>1</sup> Muthmann and Rolng, Ber., 1898, 31, 1718, cf. Mosander, Phil. May, 1843, [un.], 23, 241; Ann Chim Phys., 1844, [un.], 11, 464, Pogy Annalen, 1843, 60, 297, Marignac, Ann. Chim Phys., 1849, [un.], 27, 209; 1853, [un.], 38, 148, Watts, Q. J. Chem. Soc., 1850, 2, 140; Holzmann, J. praki Chem., 1858, 75, 321, Czudnowez, ibid., 1860, 80, 31, Bunsen, Pogy Annalen, 1875, 155, 230, 366; Hillebrand and Norton, ibid., 1875, 156, 466; Frenchs, Ber., 1874, 7, 798; Frenchs and Smith, Annalen, 1878, 191, 331.

2 Urbann, Compt. rend., 1898, 126, 835, Bull. Soc. chim., 1898, [un.], 19, 376; Ann. Chim. Phys., 1900, 1801, 120, 184.

<sup>Urbain, Compt. rend., 1898, 126, 835, Bull. Soc chem, 1898, [iii], 19, 370; Ann. Ohim. Phys., 1900, [vii], 19, 184.
James, J. Amer Chem Soc., 1908, 30, 182, 979, 1909, 31, 913; 1912, 34, 757.
Mosander, Ioc. ett., Manignac, Ioc ett.; Bunsen, Ioc. ett.; Zschiesche, J. prakt Chem., 1869, 107, 65, Brauner, Trans Chem. Soc., 1882, 41, 68; 1898, 73, 951; Monatsh., 1882, 3, 1; Ulbain, Ann. Chim Phys., 1900, [vii.], 19, 228.
Holmberg, Zeitsch. anorg. Chem., 1907, 53, 83; Bihang K. Svenska Vet-Akad. Handl., 1902-3, 28, u. No. 5; Garnier Arch. Soc., phys. nat., 1915, [vii.] 40, 93, 199.
Hiller, Incarqural Dissertation (Bellin, 1904), R. J. Meyer, Zeitch. anory. Chem., 1904, 41, 97, cf. Drossbach, Ber., 1800, 33, 3506.
Jennis and Rhodes. J. Amer. Chem. Soc., 1915, 37, 807, cf. Dennis and Bennett, 22, 1912, 24, 7.</sup> 

d., 1912, 34. 7.  $^{8}$  One hundred grams of rare earths require about 1250 grams of the carbonate solution.



B. Other Methods of Separation. - The following are among the more important of these processes

(1) Fractional Precipitation with Ammonia, 1 Sodium Hydroxide, 2 or Magnesia,3—By the gradual addition of magnesia to a boiling solution of the nitrates until the filtrate no longer exhibits the absorption spectrum of "didymum," a solution can be rapidly obtained containing lanthanum quite free from "didymum". The yield, however, is very poor.

(11 ) Auer von Welsbach's Oxide Method - By this process a fairly rapid separation of lanthanum and "didynum" is obtained. It is founded on the fact that lanthana is a stronger base than "didynua". One hundred griss. of mixed oxides are dissolved in the minimum of intric acid, and the solution diluted to 800 c c., another 100 gims of oxide are stiffed up with 100 c.c. of water and poured into the nitrate solution. Chemical change soon commences and the mass becomes warm, it is then heated on the steam-bath, with frequent shaking and stirring for some time, diluted to 1400 c.c., and allowed to stand for twelve hours. The precipitate is filtered and washed with water It contains the greater part of the "didymnun," most of the lanthamim being in the filtrate. In subsequent fractionation more than half the carths should be left as nitrates at the lanthanum end and less than half at the "didymum" end 1

(iii) Decomposition of the Nitrates.5-The mixed nitrates are mixed with excess of potassium and sodium intrates, and heated to 420°-460° until part of the rare earth intrates have been decomposed. The mass is their extracted with water, when a residue of basic "didyimum" nitrate, practically free from lanthanum, is obtained. If the aqueous extract is pink, it is evaporated to dryness and the fusion repeated, until eventually a practically colourless filtrate is obtained. Separation into cinde lantham and crude "didyima" may be stapidly effected in this manner, but pure lanthana cannot be thus obtained

(iv.) Fractional Precipitation of the Chromates.6-The oxides are dissolved in chronic acid and precipitated in dilute boiling solution with potassium chromate The order of precipitation is Ce. La. Pr. Nd, Sm

(v) Electrolysis of Aqueous Salt Solutions 7 -- When an aqueous solution of the chlorides or nitrates of the rare earth elements is electrolysed, a precipitate of mixed hydroxides appears at the cathode. By removing this precipitate from time to time, a fractionation of the rare earths in the order of their basic strengths may be effected, the weakest base precipitating first.

1713 Chem Soc, 1886, 1896, 29, 2152, Chem News, 1896, 74, 274, Brauner and Pavlíček, 1913, 331

2 Drossbach, Ber., 1896, 29, 2152, Chem News, 1896, 74, 274, Brauner and Pavlíček, 1713, Chem. Soc., 1892, 81, 1243

3 Muthmann and Rolig, Ber., 1898, 31, 1718; R. J. Meyer and Marckwald, Ber., 1900,

33, 3003.
4 Auer von Welsbach, Monatsh , 1853 4, 630; 1884, 5, 1, 508, Schottlander, Bar., 1892, 25, 378, 569

1892, 25, 378, 569
 Damour and Deville, Compt. rend., 1861, 59, 270, Bull. Soc. chim., 1861, [ii.], 2, 339; Bettendorf, Annalin., 1899, 256, 159., Schulzenberger, Compt. rend., 1895, 120, 1143; Urbain, Ann. Chim. Phys., 1909, [vii.], 19, 225., Blauner and Paylick, loc. cit.
 Bohun, Zeitsch. angew. Chem., 1902, 15, 1782
 Phenius and Lemon, J. Amer. Chem. Soc., 1915, 37, 131. Dennis, U.S.A. Pats., No. 1, 155, 33.

No. 1,115,513.

Mangnac, Ann. Chem. Phys., 1853, [nr.], 38, 148., Hermann, J. prokt. Chem., 1861,
 82, 385., Erk, Jahresber., 1870, p. 319., Cleve. Bull. Soc. chem., 1874, [nr.], 21, 196,
 246; 1883, [nr.], 39, 151, 289., 1885, [nr.], 43, 359., Branner, Mondsh., 1852, 3, 1. 486;
 Tran. Chem. Soc., 1882, 41, 68, 1883, 43, 278. Frenchs and Smith, Annalen, 1878,

This method gives a rapid separation of lanthanum from praseouymum or Middymium." It is convenient to use a solution of the nitrates contained in a tall jar, at the bottom of which a layer of mercury, kept in constant agitation by means of a stream of air, serves as the cathode, a platinum wire may be used as anote. The current should be adjusted so as to obtain a suitable rate of precipitation.

(v1) Miscellaneous.1

C. General Method of Procedure.—The separation of lanthanum, praseodymium, and neodymium is best effected in a series of stages by the fractionation of the double nitrates. The procedures recommended by Urbain and by James are practically identical 2. Thoria and ceria should have been eliminated, and also the greater part of the earths of the yttrium group The earths to be fractionated (prepared from the exalates by ignition) are dissolved in the minimum of nitric acid, an equal amount of nitric acid is neutralised by magnesia, the two solutions mixed and evaporated until small crystals form when a current of cold air is blown on the surface. A little water is sprayed on the surface, and the whole allowed to stand for twenty-four hours. An initial separation into crystals and mother liquor is thus effected. The further fractionation is then carried on, using water for solvent, as has been previously described (p. 323)

The most soluble fractions rapidly undergo change, developing a yellow colour and showing the absorption spectrum of samarium. All the eibium and yttrium salts present also concentrate into these soluble fractions The fractionation of the double magnesium salts is continued until spectroscopic examination indicates that the soluble samanum fractions are nearly free from neodymum; a result that is brought about comparatively rapidly. After fractionation has been carried out for some time, it usually happens that the final mother liquors crystallise with great difficulty, owing to the accumulation of soluble impurities in them. The solutions are then diluted, the rare earths precipitated with oxalic acid, washed, reconverted into the double magnesium intrates, and the fractionation continued. When the purified, very soluble, double nitrates again refuse to crystallise, it is due to -the accumulation of the simple intrates of the yttrium group in the final liquors. The further treatment of these soluble fractions is described later (p 349).

In a comparatively short time a practically complete separation of neodymium from samarium may be effected, and the fractions classified according to their composition into four groups, which, commencing with the least soluble, are a follows -(a) Crude lanthanum and praecodymium salts, the separation of lanthanum from praseodymium not having proceeded Every far; (\$\beta\$) praseodymium and neodymium salts (i.e. "didymium" salts); (γ) crude neodymium salt; and (δ) samarium, curopium, and gadolinium magnesium nitrates together with the simple nitrates of terbium, dysprosium,

See Marignac. Ann Chim. Phys., 1849, [iii.], 27, 209; 1853, [iii.], 38, 148; Rumnolsberg, Pogg. Annalen, 1859, 107, 631, Beilin, Inrugural Dissertation (Gottingen, 1864);
 Frighs, Ber., 1874, 7, 798; Frerichs and Smith, Annalen, 1878, 191, 331; Biauner, loc.
 Pettersson, Zeitsch anorg. Chem., 1893, 4, 1; von Scheele, stud., 1889, 17, 310;
 Ibach, Ber., 1902, 35, 2826; Biowniug. Chem. News, 1914, 110, 49; Compt. rend.,
 158, 1679; Stoddart and Hill, J. Amer. Chem. Soc., 1911, 33, 1076; Baskerville
 Turrentine. shid. 1904, 26, 46. Turentine, whid, 1904, 26, 46.

James, J. Amer Chem. Soc., 1912, 34, 757; cf. whid., 1908, 30, 979; Urbain, Chim. whis. 1908 A 57.

-holmium, yttrium. Of the rare earth elements in (δ), all but yttrium are rare in comparison with lanthanum, prascodymmum, and neodymnum; samarium and gadolimum are the most abundant.

The fractions (a) are now fractionated by Auer von Welsbach's method, for which purpose they are converted into double ammonium intrates and dissolved in water containing one-tenth their weight of intire acid. The double nitrates are then methodically fractionated, in each case, after decanting mother liquor from one fraction to the next, the crystals are washed with a little concentrated nitric acid and the washings also added to the next fraction. This is the best general procedure for the preparation of pure lanthanum and praseodymium material. Fractionation may be carried out from aqueous solution simply, but proceeds more rapidly in the presence of nitric acid 1

Each of the fractions  $(\beta)$  is converted into the double manganese intraces, and the fractionation continued from nitric acid of density 1:3. The greater part of the salt present in a fraction should be separated at each crystallisation, so that each mother liquor is small in volume compared with the volume of the crystals. This procedure is the most rapid yet known for the separation of praseodymium from neodymium, and enables the fractionation to be continued and pure salts separated at each end until the intermediate fractions are practically insignificant

The neodymnum fractions  $(\gamma)$  are fractionated still further for the preparation of pure neodymium salt, any praceodymium still present passing into the least soluble, and any sumarium, etc., into the most soluble portions. According to Demarcay, the elimination of samarium from neodymium by this method requires only a few days' work on four fractions. It is even more tapid, for the elimination of both praseodymium and samarium, to convert each of the (y) fractions into the double manganese intrates, as. with (B).

The further treatment of fractions (δ) is described later (p. 349) 2

D Purification of Lanthanum, Praseodymium, and Neodymium. --Lanthamim ammonium nitrate may be obtained in a high state of purity by prolonged fractional crystallisation, the last traces of prascodymium salt concentrating in the most soluble fractions The most rapid method for the elimination of a little prascodymnum from a nearly pure lanthanum pre-paration is said by R. J. Meyer to be the double carbonate method. The following process is recommended by James for the final purification 3

The double ammonnim intrate is dissolved in water and precipitated with oxalic acid. The oxalate is heated with concentrated sulphuric acid and The powdered thereby converted into the neutral, ambydrous sulphate salt is slowly added to ice-cold water till a saturated solution is obtained, after which it is filtered and heated to 32°. The hydrated salt which separates out is washed with hot water, dehydrated, and the crystallisation repeated. The rapid elimination of traces of prascodymium by this method

<sup>&</sup>lt;sup>1</sup> This method was the first by means of which praceodynnum and needynnum were separated. It is not, however, the best method for that purpose 
<sup>2</sup> Muthmann and Wess have separated lauthanum, praceodynnum, and needynnum very satisfactorily merely by prolonging the fractionation of the double magnesium initiates; Feit and Przibylla also recommend this process, and propose the addition of cerous magnesium intrate from time to time to the most soluble fractions, to ail in the separation (cf. p 329). James, J. Amer. Chem. Soc., 1908, 30, 979; 1912, 34, 757.

is due to the fact that lanthanum and praseodymium sulphates are not isomorphous. A pure lanthanum salt yields a pure white oxide, and when converted into the intrate or chloride and examined through a thick layer of concentrated solution exhibits no trace of the prascodymium absorption bands.

Praseodymuun material may be contaminated with lanthanum and neo-'dynnum The last traces of lanthanum are best removed by a process in which they accumulate in the most soluble fractions, e.g. crystallisation of the ethylsulphates, bromates, or the simple intrates. Crystallisation of the oxalates from nitric acid is also a very expeditious method. According to Brauner,2 pure prascodymium tetrovide, Pr<sub>2</sub>O<sub>0</sub>, separates when the nitrate is fused with nitre at 400°. The last traces of neodymium may be removed by the prolonged crystallisation of the double animonium, magnesium or manganese nitrate, or by the double carbonate method (p. 312).

According to Baskerville and Turrentine, pure praseodymium material is readily obtained in one operation from an impure material containing not more than 10 per cent of lanthanum The hydroxides are precipitated with ammonia, thoroughly washed till free from ammonia, dissolved in an ice-cold, saturated solution of citric acid, and the solution filtered and heated to 100° Normal prascodymium citrate separates as an amorphous gicen powder, which is filtered off and washed with hot water.3

Neodymium material is hable to contain praseodymium and samarium. The latter, however, may be eliminated very easily by the crystallisation of the double ammonium, magnesium, or, best of all, manganese nitrates complete elimination of praseodymium is a very tedious operation, one of the best methods consists in the fractional crystallisation of the nitrate from concentrated nitric acid, which concentrates the prascodymium salt in the most soluble fractions 4 Crystallisation of the bromate likewise climinates the praseodymium in the most soluble fractions Baskerville and Stevenson recommend fractional precipitation of the chloride from acid solution (by saturation with hydrogen chloride) as a ready means of eliminating lanthanum and praseodymium chlorides 5, while Holinberg and Garmer purify neodymium material by fractionating the metanitrobenzoate. Strongly ignited neodymium oxide has a beautiful blue colour, several tenths of one per cent of praseodymia, however, do not affect the colour The absorption bands in the blue are sharply defined when a neodymium intrate or chloride solution is free from samarium and prascodymium salt, but are somewhat hazy in the presence of these impurities.

SEPARATION OF SAMARIUM, EUROPIUM, GADOLINIUM, AND TERBIUM.

The double potassium (sodium) sulphates of these elements are slightly, but appreciably, soluble in a saturated solution of potassium (sodium) sulphate, and hence divide themselves between the precipitate and the filtrate when

<sup>1</sup> From intric acid; Feit and Przibylla, Zeitsch. anorg Chem, 1905, 43, 202.
2 Brauner, Proc. Chem. Soc., 1898, 14, 70, 1901, 17, 66.
3 Baskerville and Turnentine, J. Amer. Chem. Soc., 1904, 26, 46; Baskerville, Zeitsch. anorg. Chem., 1905, 45, 86. The method is adversely criticised by R. J. Meyer (ibid., 1904, 41, 97) For another method of eliminating La, see Orloff, Chem. Zeit, 507, 31, 115
307, 31, 115
4 Baskerville and Stevenson, J. Amer. Chem. Soc., 1911, 33, 1.
8 Baskerville and Stevenson, J. Amer. Chem. Soc., 1904, 26, 54.

the potassium (sodium) sulphate method is used to effect a separation of the cerium from the yttrium earths. If the sulphate method be applied in such a manner as to separate the cerum double salts in a series of fractions, samarium, europium, gadolimum, and terbium may be concentrated in the final fractions - It was by operating in this manner that Lecoq de Boisbaudran, the discoverer of samarium, was able to concentrate samarium with "didymium," and afterwards separate them by various processes and a similar method of experimenting led Marignac to the discovery and isolation of gadolinium.1

The methods which have been applied to effect the separation of samarium, europium, gadolinium, and terbium from one another or from the elements of the cerum and yttrum groups are as follows

- A. Fractional Crystallisation, -The following salts have been

(i) Formates,  $^2$  M(O.OC H),  $_3$  HI<sub>2</sub>O, from water (ii) Oxalates,  $^3$  M<sub>2</sub>(C<sub>2</sub>O<sub>4</sub>),  $_3$  eH<sub>2</sub>O, from dilute intric acid (iii) Ethylsulphates,  $^4$  M(C<sub>2</sub>H<sub>5</sub> SO<sub>4</sub>), 9H<sub>2</sub>O, from water (p. 340).

(iv) Nitrates, M(NO<sub>3</sub>), 511,0, from nitric acid of density 1.3

- (v.) Double magnesium intrates, 2M(NO<sub>3</sub>)<sub>3</sub> 3Mg(NO<sub>3</sub>)<sub>2</sub> 24H<sub>2</sub>O<sub>3</sub> from nitric acid (13)
- (vi.) Double manganese intrates,7 2M(NO<sub>3</sub>), 3Mn(NO<sub>3</sub>), 24H<sub>2</sub>O<sub>5</sub> from mtric acid (1 3)
- (vn.) Double nickel intrates, 8 2M(NO<sub>3</sub>)<sub>2</sub> 3Ni(NO<sub>3</sub>)<sub>2</sub> 24H<sub>2</sub>O<sub>3</sub>, from intric acid (13)

(vii) Picrates,  $^{0}$  M[O  $C_{6}H_{2}(NO_{2})_{3}]_{3}$   $H_{2}O_{5}$  from water. (ix) Metanitrobenzenesulphonates,  $^{10}$  M[ $C_{6}H_{4}(NO_{2})$  SO<sub>4</sub>] $_{4}$   $_{2}H_{2}O_{5}$  from

Lecoq de Borsbaudian, Compt. rend., 1879, 88, 322, 1879, 89, 212; Mangnac, Arch. Sc. phys. nat., 1880, [iu.], 3, 413, Compt. rend., 1880, 90, 899., Ann. Chim. Phys., 1880, [v.], 20, 535.
 Belafontaine, Arch. Sc. phus. nat., 1877, 59, 176, 1878, 61, 273., Ann. Chim. Phys., 1878, [v.], 14, 238.
 Mangnac, los. cet., Kiuss and Hofmann, Zeitsch. anory. Chem., 1893, 4, 27., Fett, ibid., 1905, 43, 267., Bettendorf, Annalen., 1907, 352, 88., Urbain, Ann. Chim. Phys., 1900, [vii.], 19, 194.
 Belafontaine, Arch. Sci. phys. nat., 1865, 22, 30, 38, 1878, 61, 273; Ann. Chim. Phys., 1878, [v.], 14, 238.
 Mangnac, Arch. Sci. phys. nat., 1878, 61, 273; Ann. Chim. Phys., 1878, [v.], 14, 247.
 Urbain, Ball. Soc. chim., 1898, [iii.], 19, 376., Compt. rend., 1898, 126, 835., 1904, 139, 736., 1906, 142, 785., Ann. Chim. Phys., 1900, [vii.], 19, 184., J. Chim., phys., 1906, 4, 334.
 Demarçay. Compt. rend., 1896, 122–728, 1900, 130, 1019., Benedicks. Zeitsch. anorg. Chem., 1900, 22, 393; Urbain, Compt. rend., 1904, 139, 736; 1900, 149, 37., Feit and Przibylla, Zeitsch. anorg. Chem., 1900, 22, 393; Urbain, Compt. rend., 1904, 139, 736; 1900, 149, 37., Feit and Przibylla, Zeitsch. anorg. Chem., 1909, 130, 1019, 1185, 1469, 131, 313, 1901, 132, 1484., Urbain and Lacombe, abid., 1903, 137, 792., 1901, 138, 84, 627. 1166. Urbain, J. Chim., phys., 1906, 4, 31, 105., Vulthiman and Wess, Annalen., 1904, 31.; Feit and Przibylla, loc. cit.; James and Robinson, J. Amer. Chem. Soc., 1911, 33, 1363, Holmberg, vide infra.
 Lacombe Rull. Soc. chim., 1904, 1iu.], 21, 570; Chem. News. 1904, 80, 277.

vide infra.

7 Lacombe, Bull Soc. chim, 1904, [iii.], 31, 570; Chem. Neus, 1904, 89, 277.

8 Urbain, Compt. rend, 1904, 139, 736, 1905, 141, 521; J. Chim. phys., 1906, 4, 824.

9 Holmberg, Bihang K. Svenska Vet Akad Handle, 1902-3, 28, 11. No 5, Zerlsch.

1 Remostt J. Amer. Chem. Soc., 1912, 34, 7; Dennis

anory Chem., 1907, 53, 83, Dennis and Bennett, J. Amer. Chem. Soc., 1912, 34, 7; Dennis and Rhodes, 1911, 1915, 37, 807.

10 Holmberg, loc. cit.

(x.) Bromates, M(BrO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O, from water. (xi) Dimethylphosphates, M[(CII<sub>7</sub>)<sub>2</sub>PO<sub>4</sub>]<sub>3</sub> xH<sub>2</sub>O, from water.

(xii ) Chlordes, MCl, 6H2O, from hydrochloric acid

The solubilities of corresponding salts of samarium, europium, gadolinium and terbium usually increase in the order Sm, Eu, Gd, Tb limethylphosphates the reverse holds good; with the simple nitrates the solubilities diminish in the order Sm, Eu, Gd, but the solubility of the terbium salt exceeds that of the gadolinum. In fact, from gadolinum to lutecium and celtium the "serial order" holds good (see p 327).

B. Fractional Precipitation.—The following compounds may be

fractionally precipitated:

\$0 . · ·

(i.) The double sulphates.4

(ii.) The hydroxides or basic salts, by ammonia.5

(in.) The stearates.

The hydroxides of terbium, samarum, and gadolinium are precipitated in the order Tb, Sm, Gd by ammonia, europium hydroxide piecipitates before the hydroxide of samarium

C. General Method of Procedure.-When the earths extracted from a muneral contain very little of the yttrium group and the fractionation is commenced by the double magnesium nitrate method without previously eliminating the jttrium group, the whole of the samarium, curopium, gadolinium, etc., finds its way into the soluble fractions ( $\delta$ ) previously mentioned (p. 344). If, however, the earths extracted from a numeral have been untially submitted to the double sulphate separation process, the crude double sulphates of the cerium group contain probably the greater part, but not all of the samarium, europium, and gadolinium, and very little of the terbium, while the filtrate contains the remainder of these elements as sulphates. From a mixture of crude yttrium earths the elements samarium, europium, gadolinium, and terbium may be eliminated in the least soluble fractions by fractionally crystallising the ethylsulphates or bromates, as has been previously mentioned (p 340)

In outlining the procedures best adapted to the separation of samarium, europium, gadolinium, and terbium, it is only necessary to describe the method of working up the extremely soluble fractions ( $\delta$ ) from the double magnesium nitrate fractionation of the cerum group (p 311) fractions obtained from crude yttrium earths by the ethylsulphate or bromate method may be converted into double magnesium nitrates and treated in

a similar manner.7

# HE RABE EARTH ELEMENTS.

The fractionation of the double magnesium nitrates (8) is continued from nitric acid of density 1.3, as recommended by Demarçay. This is the best method for separating samarium, curopium, and gadolimium. The fractionation should be continued until a considerable quantity of the samarium has been eliminated in the head fractions, free from curopium 1 Throughout the course of this fractionation great care must be taken to prevent the deposition of crystals of magnesium nitrate, which are apt to separate out, since that salt is less soluble than the double intrates. The solutions should therefore be allowed to cool down and form supersaturated solutions, and then seeded either with tiny crystals saved for the purpose, or, better, with bismuth magnesium nitiate 2

After as much europium-free samarium as possible has been removed, considerable quantities of bemuth magnesium nitrate are continually added to the final mother liquors as the fractionation proceeds. The material should be separated into about 20 fractions and fractionation carried on daily for about six months. After each crystallisation the volume of the mother liquor should be about one-tenth of the total volume. By working in this manner on 611 grams of rare earths, which consisted almost entirely of samana and gadolinia, Urbain and Lacombe (p. 327), the originators of this method, added in all about 5 kilos of bismuth magnesium mitrate. They eliminated the head fractions as soon as it was clear that they were free from curopium. Finally, after nearly six months' fractionation, the two head fractions were found to be practically devoid of samarium and europium, i.e. they were practically pure bismuth magnesium intrate. The remaining fractions contained much bismuth salt and the following amounts of rare earths (head fraction was No 12) -

19 20 21 22 15 16 17 18 No. of fraction . 11 0.537 1.01 1.50 1.74 1.50 1.80 2.7 12 7 29.0 grams. Wt of oxides

Numbers 14 to 19 were free from gadolimum, and numbers 23 onwards were free from europium. Thus the earths extracted from fractions 14 to 19 consisted of pure europia, and the intermediate fractions between pure curopia and pure gadoluna had been reduced to three (numbers 20, 21, and 22) 8

<sup>1</sup> By examining the absorption spectrum of the double intrates, inclied in their water

<sup>1</sup> By examining the absorption spectrum of the double intrates, melted in their water of crystallisation, through a thickness of about 12 cms (the diameter of a litte flash), and rejecting fractions which do not show the two bands 5337 and 5251 due to europium, considerable amounts of samarium but practically no europium may be chimiated.

2 It has been already mentioned (see p. 344) that considerable trouble may be experienced in crystallising the most soluble part of the (8) fractions owing to the first that all the yttrium earths left in the clude cerium carths become concentrated in them. The following means are available for climinating the yttrium earth from these uncrystallisable fractions.—

(1) Convert into sulphates and apply the double sulphate method.

(ii) Convert into the sumple nitrates and fractionally crystallise from intric acid of density 1.3 This excellent method was strongly recommended by Demarcay (Compt. rend., 1900, 130, 1019), and has also been employed by James (J. Amer. Chem. Soc. 1913, 25, 235). The intric acid should not have a greater density than 1 3, or yttrium intrate timydrate,

The intro said should not have a greater density than 13, or yttrium intrate trihydrate, Y(NO<sub>3</sub>), 3H<sub>2</sub>O, may separate and cause trouble

(iii) Add a large quantity of bismuth magnesium nitrate and fractionally crystallise from intric acid (Urbain and Lacombe, Compt. rend., 1904, 138, 84, J. Chim. phys., 1906,

<sup>4, 121)

3</sup> Without the addition of the isomorphous bismuth salt, the separation of suropium from the separation of the isomorphous bismuth salt, the separation of suropium from the separation of the isomorphous bismuth salt, the separation of suropium from the separation of the isomorphous bismuth salt, the separation of suropium from the separation of the isomorphous bismuth salt, the separation of suropium from the separation of the isomorphous bismuth salt, the separation of suropium from the separation of the isomorphous bismuth salt, the separation of suropium from the separation of the isomorphous bismuth salt, the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separation of suropium from the separatio gadolinium requires 18 intermediate fractions (Demarkay, see p 347). The work of Urbain and Laconibe has been repeated and vented by James and Robinson (p 347), who worked on a larger scale than the French chemists.

From the preceding description it will be seen that the separation of samarium and europium by Urbain and Lacombe's method is quantitative. For the purification of samarium it is necessary to climinate the last traces of neodymum. This may be accomplished (i) by the fractional crystallisation of a mixture of equal weights of the double nitrate of samarium and magnesium and the double intrate of bismuth and magnesium, when the neodymium is eventually found in the head fraction, 1 (ii ) by the fractional crystallisation of samarium dimethylphosphate, when the neodymium is rapidly expelled in the most soluble fractions, 2 or (m.) by the fractional crystallisation of samarium manganese m<br/>trate from m<br/>tric acid.  $^3$ 

The clumnation of terbium from gadolinium is more tedious than the elimination of europium. The fractions following those which contain the europium are suitably treated to extract the iare earths and to convert them into the simple intrates. The intrates are then fractionated from nitile acid of density 1.3 in the presence of a large excess of bismuth nitrate,  ${\rm Bi(NO_3)_3~5H_2O}$ . The order of separation is Gd, Bi, Tb, (Dy), and a practically quantitative separation of gadolmium and terbium can be brought about by prolonging the fractionation for a sufficient length of time. The separation proceeds very slowly, but it is certain

The purification of terbium is completed when both gadolinium and dysprosium have been eliminated from it. The climination of the gadolinium has just been described; at the same time that the separation of gadolinium and terbium is proceeding in the least soluble fractions, a very good, but not quantitative, separation of terbium and dysprosium simultaneously takes place in the more soluble fractions, and, owing to the presence of the excess of bismuth salt, the intermediate terbium-dysprosium fractions are reduced to a very small amount

The great objection to the method of the fractional crystallisation of the nitrates for separating gadolinium, terbium, and dysprosium is the extreme slowness with which it effects the separation A much more rapid and convenient process is the fractional crystallisation of the bromates, which separates gadolinium and terbium particularly well. Moreover, it is very probable that the separation can be conducted quantitatively in the presence of neodymum bromate.

The best alternatives to the methods outlined in the preceding paragraphs for the isolation of gadolinium and terbium are. (i.) the fractionation of the double nickel nitrates to eliminate terbium from gadolinium, and vice versa; (ii.) the fractional precipitation of the hydroxides with ammonia to climinate the last traces of gadolinium from terbium prepared by method (i.); (iii) the fractionation of the ethylsulphates to eliminate dysprosium from terbium, 4 and (iv.) the fractionation of the dimethylphosphates, which rapidly eliminates terbium from gadolinium.

#### THE SEPARATION OF THE YTTRIUM GROUP.

The metals of this group are distinguished from those of the cerium and "terbium groups by the solubility of their double potassium sulphates in a

<sup>&</sup>lt;sup>1</sup> Urbain and Lacombe, Compt. rend., 1904, 138, 1166; Urbain, J. Chim. phys., 1906, 4,

J. C. Morgan and C. James, J. Amer. Chem. Soc., 1914, 36, 10.
 Lacombe, Bull. Soc. chira., 1904, [iii.], 31, 570, Chem. News, 1904, 89, 277
 For references, see p. 351.

saturated solution of potassium sulphate, and by the inability of their nitrates to form double salts with the alkali nitrates or the nitrates of magnesium, zinc, nickel, etc.

The separation of the elements of the yttimm group from one another has, as yet, not been effected so satisfactorily as the separation of the elements of the other groups. Practically all the methods that have been tried are mentioned in the following paragraphs, and an outline of the experimental procedure is given in the cases of the more valuable methods

A Fractional Crystallisation .-- The following salts have been employed -

(1.) The ethylsulphates, from water 1 The utility of this valuable method is discussed later (pp. 355-6).

(n ) The bromates, from water.2 The utility of this method is also discussed later (pp 355-6)

(111) The acetates, from acetic acid.3

(iv ) The metantiobenzoates, from water 4

(v.) The nitrates, from strong mitric acid. The utility of this valuable method is discussed later (pp 356-7)

(vi.) The double ammonium oxalates . The wet oxalates are dissolved in a boiling, concentrated, slightly ammoniacal solution of ammonium oxalate, and the solution cooled in successive stages, the crystals separating at each stage being collected before proceeding to the next.7

(vii ) The acctylacetonates, from alcohol and benzene 8 (viii ) The piciates, from water a. This is an efficient method for removing small amounts of the earths of the cibium and yttium groups from those of the certuin group and for separating yttrium from erbium and holmium.

(iv) The chlorides, from concentrated hydrochloric acid. 10

71, 226.

5 lemargay, Compt. rend , 1896, 122 728 , 1900, 130, 1019 , Urbain, ibid , 1907, 145, 759 , 1908, 146, 406 , Bull Soc chim , 1905, [in ], 33, 739 , Blumenfeld and Urbain, Compt rend , 1914, 159, 323 , Holmberg, loc cit.

6 Auer von Welsbath, Monath , 1906, 27, 935 , 1905, 29, 181 , 1911, 32, 373 , Chem. News, 1907, 95, 197 , 1908, 98, 223 , Holmberg, loc cit , Holmann and Burger, Ber , 1908, 47, 308.

7 For other methods of fractionating by means of oxalates, see Mosander, Phil. Mag., 1843, [iii.], 23, 241, Delafontaine, Arch Sci phys nat., 1804, 21, 97, Annalen, 1864, 134, 99, Cleve, Compt rend, 1879, 89, 419, Lecoy de Boisbaudian, ibid, 1879, 88, 322; 1886, 103, 627; Kruss, Annalen, 1891, 265, 1; Ubain, Ann. Chim. Phys., 1900, [vii.], 19, 184, Marc, Br., 1902, 35, 2:82, Dennis and Dales, J. Amer. Chem. Soc., 1902, 24, 401.

\*\* Urbain and Budischovsky, Compt. rend., 1897, 124, 618, Urbain, Ann. Chim. Phys., 1900, [vii.], 25, 124.

1900, [vi ] 19, 184.

9 Holmberg, Zeitsch. anorg Chem., 1907, 53, 83; Dennis and Bennott, J. Amer. Chem. Soc., 1912, 34, 7, Dennis and Rhodes, ibid., 1915, 37, 807, James, ibid., 1911, 33, 1832.

10 Dennis and Dales, J. Amer. Chem. Soc., 1902, 24, 401; Langlet, Arkiv Kem Min. Geol., 1907, 2, No. 32, James and Bissel, J. Amer. Chem. Soc., 1914, 36, 2065.

Utbain, Bull Now thim, 1898, [in], 19, 376, Compt. rend., 1898, 126, 835, 1906,
 142, 785, Ann Chim. Phor., 1900, [vn], 19, 184, G and E Utbain, Compt. rend., 1901,
 132, 136, R. J. Meyer and Wuomen, Zeitsch anny Chem., 1913, 80, 7.
 James, J. Amer. Chem. Soc., 1908, 30, 182, 979, 1910, 32, 517; 1911, 33, 1332;
 1912, 34, 777, 1913, 35, 235, Holden and James, thut, 1914, 36, 638, James and Bissel, thut, 1914, 36, 2005. In the first of these papers, James mentions that the fractionation of the bromater is superior to the fractionation of any of the following salts—Sulphtes, necessary, authorize, methyleadistics nearly supplying converges refered by superiors. succinates, xanthates, methylsulphates, propylsulphates, camphorates, indates, thoo yanates, monochloracetates, monochloracetates, monochloracetates, monochloracetates, monochloracetates, monochloracetates, monochloracetates, monochloracetates, fungue of Dissertation (Munich Tech Inst., 1902)

4 Holmberg, Arkiv Kem Min Giol., 1911, 4, Nos 2, 10, Zeduh anory Chem., 1911,

(x.) The dimethylphosphates, from water.1 This separation appears to . very rapid one, and deserves further study.

B. Fractional Precipitation. — The following compounds may be

ractionally precipitated :-

(1) The hydroxides or basic salts, by the addition of ammonia, sodium hydroxide, aniline, or magnesia. These processes are very tedious. The nagnesia method is useful in removing small amounts of impurities from ottruum, for which purpose the magnesia is stured up into a "cream" with water, and added gradually to a concentrated, boiling solution of the rare earth nitrates

(11.) The chromates 6 For this purpose the rare earths (i.e. the oxides) are mixed with the calculated amount of chromic acid, and water then added A violent reaction cusues and an aqueous solution of the dichromates is produced.7 The liquid is diluted (about 1 litre per 50 grams of oxides) and 10 per cent potassium chromato added until a permanent turbidity is produced The liquid is heated to boiling, and while heating by a burner is still carried on, a vigorous current of steam is blown through the liquid. By this means a constant volume may be maintained throughout the fractionation, which is effected by adding 10 per cent. potassium chromate, drop by drop, from a tap-funnel. The precipitate is collected in a series of fractions

The method is an excellent one for the rapid preparation of practically pure yttrium material from a starting material composed of the crude yttrium earths from which the cerium group, and preferably the terbium group, has been eliminated. Unfortunately it does not seem to have any other practical use.

(iii.) The ferrocyanules 8 This method, like the preceding, enables pure

yttrium material to be obtained, but the yields are very poor.

(iv.) The succenates.9 A solution of ammonium or sodium succenate is added, drop by drop, to a 2 or 3 per cent boiling solution of the nitrates, and the precipitate collected in fractions.

(v.) The stearates. 10 A 50 per cent. hot alcoholic solution of potassium

1 J. C. Morgan and C. James, J. Amer. Chem. Soc., 1911, 36, 10.

2 Mosander, Phil. Mag., 1813, [iu.], 23, 251., Lecoq de Borsbaudran, Compt. rend., 1886, 102, 1003, 1005, Kruss. Annalen, 1891, 265, 1. Muthmann and Bam, Ber., 1900, 33, 1743; Marc. sbid., 1902, 35, 2782; Urbam, Ann. Chim. Phis., 1900, [vii.], 19, 184. G. and E. Urbam, Loc. ett.; Postius, loc. ett., Dennis and Dales, loc. ett., Bettendorf, Annalen, 1907, 352, 88, Holmberg, loc. ett., Dennis and Dales, loc. ett., Bettendorf, Annalen, 1907, 352, 88, Holmberg, loc. ett.

3 Holden and James, J. Amer. Chem. Soc., 1911, 36, 638.

4 Kruss, Annalen, 1891, 265, 1. Zeitsch. anorg. Chem., 1893, 3, 108, 353; K. A. Hofmann and Kuss. shid., 1893, 3, 407, 4, 27

5 Muthmann and Rolig, Ber., 1898, 31, 1718; Drossbach, Ber., 1896, 29, 2452; Dennis and Dales, loc. cit., James, J. Amer. Chem. Noc., 1907, 29, 495

and Dales, loc. cit., James, J. Amer. Chem. Noc., 1907, 29, 495

c. Kruss and Loose, Zeitsch. anorg. Chem., 1893, 3, 92, Morssan and Etard, Compt., etc., 1894, 122, 573; Muthmann and Bohm, Ber., 1900, 33, 42, Bohm, Zeitsch. angew. Pend., 1894, 122, 573; Muthmann and Bohm, Ber., 1900, 33, 42, Bohm, Zeitsch. angew. Chem., 1902, 15, 1222; Dennis and Dales, loc. cit., R. J. Meyer and Wuotunen, Zeitsch. anorg. Chem., 1913, 80, 7; Egan and Balke, J. Amer. Chem. Soc., 1913, 35, 365, Holden and James, while, 1914, 36, 638.

7 Instead of using a solution of the dichiomates, a neutral solution of the intrates may be used and potassum dichromate added (James, J. Amer. Chem. Soc., 1912, 34, 769).

8 Rowland, Johns Hopkins Univ. Circular, 1894, 112, 73; Chem. News, 1894, 70, 68; Jones, Amer. Chem. J., 1895, 17, 154; Crookes, Chem. News, 1894, 70, 81; Bettendorf, Annalen, 1907, 322, 88; R. J. Meyer and Wuornen, loc. cit.

10 Lenher, J. Amer. Chem. Soc., 1908, 30, 572; Benner, while, 1911, 33, 50.

stearate is added, drop by drop, to a cold, vigorously stirred solution of the neutral intrates, and the precipitate removed in fractions. This is said to be a rapid method for preparing pure yttiia

(vi) The volates. A solution of potassium iodate is added, drop by drop, to a hot solution of the nitrates, slightly acidified with intric acid. The method is of value in the purification of yttirum, which separates last

(vn) The cobalticyanides? This affords a rapid method for the separation of pure yttrum

(viii) The tartrates A solution of tartaric acid in acetone is added to one of the rare earth nitrates in the same solvent, and the precipitate removed in

(iv ) Other salts. The acides, hypophosphites, phosphites, phosphates, arsenates, sulphites, timpstates, superioristics, phosphates, phosphates, phosphates, glycollates, mono- and di-methylphosphates, pheno yacetates, camphorates, m-nitrobanzontes, and a obenzenesulphonates have all been examined.

Other Methods.—These include the following

(1) Fractional Decomposition of the Nitrates by Heat."-For this purpose, the hydroxides are dissolved in intiic acid, the solution evaporated to dryness, and the residual nitrates heated. The evolution of red fumes is allowed to proceed until the surface acquires a peculiarly steely appearance, but should not be allowed to continue until the mass becomes creamy The fused mass is then poured very carefully into cold water, and the liquid heated to boiling. If y'trium predominates very largely over the other rare earth metals present, the entire mass generally passes into solution, and, on cooling, basic intrates separate out in the crystalline form, otherwise an insoluble residue is left, in which case it must be well agitated by vigorous boiling before the liquid is cooled. After cooling, the residue is separated by filtration, dissolved in nitric acid, and again partially decomposed, while the filtrate is evaporated to dryness, and the residue also subjected to the same treatment, and so on, the filtrate from one fraction being combined with the basic salts separated from the next

This is one of the classic methods for the fractionation of the rare It has been of great service to past investigators, scandium, ear ths ytterbium, thulium, and holinium having, in fact, been discovered as the result of the application of this method to the fractionation of the yttrium earths. At the present time it still remains one of the best methods, at least on a large scale, for eliminating erbinin from vttrum

<sup>&</sup>lt;sup>1</sup> R. J. Meyer and Wuormen, Zeitsch. among. Chem., 1913, 80, 7, Holden and James,

J. Amer. Chem. Soc., 1914, 36, 638

<sup>2</sup> Bonardi and James, J. Amer. Chem. Soc., 1915, 37, 2612

<sup>3</sup> Bernebey, J. Amer. Chem. Soc., 1912, 34, 1174.

Whittemore and James, J. Amer. Chem. Soc., 1918, 35, 127. Holden and James, loc. cit.

Jantsch and Grunkraut, Zeitsch anorg Chem , 1912, 79, 305.

Jantsch and Gunktaut, Zeituh anong Chem., 1912, 79, 305.
 Dennis and Dales, J. Amer. Chem. Soc., 1902, 24, 401
 Holden and James, J. Amer. Chem. Soc., 1911, 36, 638
 The older references are Beilin, Ferhandt Stand Naturf Kjobenhava, 1860, 8, 448,
 Bahi and Bunsen, Anaden, 1866, 137, 1, Cleve and Hogland, Bihang K. Svenska Vet. Akad. Handl., 1873, 1, No. 8, Bull. Soc. chim., 1873, [n], 18, 198, 289, Marignac, Arch. Soc., phys. nat., 1878, [n], 64, 97. Compt. rend., 1878, 87, 578, Ann. Chim. Phys., 1878, [v.], 14, 247, Nilson, Compt. rend., 1879, 88, 645, Cleve, thid., 1880, 91, 381
 Modern references are G. and E. Urbain, Compt. rend., 1901, 132, 136; Dennis and Dales, J. Amer. Chem. Soc., 1902, 24, 401, James and Pratt, thid., 1910, 32, 87, Holden and James, thid., 1914, 36, 638. See also Barnebey, loc. cit.
 VOL. IV.

Before applying this method to the fractionation of the yttrium earths, it is important the curopium, gadolinium, terbium, dysprosium, and holmium should as far as possible be eliminated (and, of course, the cerium group). Serious errors have been made owing to the fact that this initial step was not taken (see pp. 355, 363).

(11) Aver von Welsbach's Oxide Method 1 -- The manner in which this

method is carried out has been already described (p. 343)

(iii.) Hydrolysis of the Phthalates.2 -- A cold solution of the phthalates is slowly heated and the precipitated basic salts removed in fractions The

yttrium precipitates last

(IV) The Ammonium Carbonate and Acetic Acid Method 3-The hydroxides of the rare earth elements of the yttrum group are soluble in ammonium carbonate. Fractionation may be effected by dissolving the hydroxides in ammonium carbonate, adding concentrated acetic acid till the liquid becomes turbid, and then adding dilute acetic acid, drop by drop, to the well-stirred liquid until the earths are partially precipitated.

Torbium and yttrium rapidly concentrate in the first fractions, erbium and "ytterbium" in the last, and holinium, thuhum, and dysprosium in the

middle fractions

(v) The Oxalate-Carbonate Method 5-The oxalates are dissolved in a warm saturated solution of normal ammonium carbonate and then heated to 100°. The precipitate that forms is collected in fractions. The elements separate in the following order. (Tb, Y), (Dy, He), E1, Tin, (Yb, Lu) The separation of erbium from holmium, terbium, and dysprosium is very rapid

(vi.) The Basic Nitrite Method. - A neutral solution of the nitrites is boiled and stirred by a current of steam, while concentrated sodium mitrite solution is slowly added to the liquid. The precipitate of basic mitrites (!) is

removed in fractions as usual.

The separation of all the members of the yttimm group appears to be effected by the method, but at present it has only been studied in any detail so far as the separation of yttrium and eibium is concerned. For this separation it is the most rapid method known Unfortunately, however, the precipitates are at times obtained in a colloidal form, and when this occurs no appreciable separation can be effected. The formation of the colloidal precipitates is connected with the presence of excess of sodium intrate, but the necessary conditions for successful fractionation still require to be discovered

(vn.) Fractional Sublimation of the Chlorides 7 - This method is of use in separating small quantities of lutecium from ytterbium (neoytterbium), the lutecium chloride being the more volatile. Scandium and thorium are liable to be present when crude "ytterbium" is isolated, their chlorides are much more volatile than lutecium chloride

Auer von Welsbach, Monatsh, 1883, 4, 630; 1906, 27, 935, Chem. News, 1907, 95,
 197; Bettendorf, Annalen, 1907, 352, 88; cf. Drossbach, Rev., 1902, 35, 2826
 R. J. Meyer and Wuormen, Zeitsch. amony Chem., 1913, 80, 7.
 Bennis and Dales, J. Amer. Chem. Soc., 1902, 24, 401.
 Mosander, Annalen, 1813, 48, 219; Phil. May, 1843, [un.], 23, 251; Kruss, Annalen, 1891, **265**, 1

James, J. Amer. Chem. Soc., 1907, 29, 495. Chem. News, 1907, 95, 181.
 Holden and James, J. Amer. Chem. Soc., 1914, 36, 1418.
 Urbain, Bourion, and Maillard, Compt. rend., 1909, 149, 127., cf. Pettersson, Zeitsch. anorg. Chem., 1893, 4, 1.

(viii ) Electrolysis of Aqueous Salt Solutions.1-The procedure necessary has been already described (see p 313). By this method yttim may be separated from the erbia earths with considerable rapidity

(ix) Fractional Decomposition of the Sulphates by Heat?

D General Procedure. - In nearly all cases in which the yttrium group is separated by fractional crystallisation, the elements separate in serial order," with dysprosium in the least and Intecium and celtium in the most soluble fractions. The reverse order of solubilities, however, holds for the dimethylphosphates

Separation methods which depend on differences of basicity, particularly the precipitation with ammonia, decomposition of the nitrates, and crystallisation of the ovalates from intric acid, received considerable attention in the early days of rare earth chemistry. In all such methods as these, yttria acts as the strongest base present, erbia, thillia, ytterbia, and litteria as much weaker bases, and dysprosia and holima as bases interinculate in strength, Accordingly, by applying method (C 1), the fractional decomposition of the mitrates by heat, yttria may be largely separated from the other carths is separated, however, in an impine state, being contaminated more especially with holina and dysprosia, and also with terbia and gadolinia if these earths are present. Impure yttra thus obtained from a mixture of yttrium earths resists fractionation in a very extraordinary fashion, so much so that in 1896 Schutzenberger and Boudonard, as the result of their experiments on the nitrate method, were inclined to believe that among the elements of the yttrium group there is one, of atomic weight 97, that had previously been overlooked. The oxide of this new "element, 'however, as Urbain showed in 1900, was merely yttria contaminated with terbia, gadolinia dysprosia, and holma, in spite of the fact that it could not be fractionated by the partial decomposition of the nitrates, or the tractional crystallisation of the sulphates, acetylacetonates, etc.

From the preceding remarks it will be seen that it is desirable to eliminate the terbum group, dysprosium and holmium, before endeavouring to fractionate the remaining earths of the yttrimin group. It is therefore the best plan to commence by fractional crystallisation, and for this purpose the ethylsulphates or bromates should be selected. The method of procedure in the case of the ethylsulphates has been already described (p. 310), and the fractionation of the bromates may be carried out in a similar manner ethylsulphate and bromate methods may to a certain extent be looked upon as effecting a separation into three sub groups (a) the least soluble fractions, in which dysprosium, holimum, and yttimm separate in the order named (β) the middle fractions, constituting the greater part of the material and consisting essentially of yttrium salt, contaminated chiefly with dysprosium and holmium in the least soluble, and with erbium in the most soluble, fractions; and (y) the most soluble fractions, in which erbium, thulium,

<sup>&</sup>lt;sup>1</sup> Kiuss, Zeitsch anorg Chem., 1893, 3, 60. Dennis and Dales, J. Amer. Chem. Soc., 1902, 24, 401, Dennis and Lemon, dad., 1915, 37, 131; Dennis and van der Meulen, bid., 1915, 37, 1963, Dennis, U.S., I. Pat., No., 1,115,513

<sup>2</sup> Hofmann and Burger, Ber., 1908, 41, 308

<sup>3</sup> Schutzenberger and Bondounal, Compt. rend., 1896, 122, 597, 1896, 123, 782, Bull. Soc. chem., 1898, [11], 19, 227, 236, Bondouard, Compt. rend., 1898, 126, 1618. Bull. Soc. chem., 1898, [11], 19, 603, Urbain and Budischovsky, Compt. rend., 1897, 124, 618, Bull. Soc. chem., 1897, [11], 17, 98, Urbain, Ann. Chim. Phys., 4900, [vii.], 19, 184, Drossbuch, Ber., 1896, 29, 2452.

yttrium, and lutecium separate in the order named. The terbium group fractions, if any, precede the (a) fractions.

The (a) fractions serve for the preparation of dysprosium and holimum. To isolate dysprosium in a state of purity it is necessary to prolong the fractional crystallisation of the ethylsulphates containing dyspiosium until the degree of hydrolysis becomes too great for further progress to be made. By this means terbum may be satisfactorily eliminated in the least soluble fractions For further purification each fraction may be separately converted into the intrate and the fractionation continued from concentrated intric acid. This method allows a very slow but sure separation of dysprosium from holmium to be accomplished 1. At the same time it eliminates terbium, if present, in the least soluble fractions, and this may be greatly facilitated if a large quantity of bismuth intrate is added (see p. 328)

The separation of holmium from both dysprosium and yttimin is an extremely difficult task. Urbain found that the fractional crystallisation of the cthy sulphates enables erbrum, thulum, etc., to be separated from holmnum very satisfactorily, and also brings about a considerable separation between holmmin and dysprosium, and that the subsequent crystallisation of the nitrates brings about the climination of the remaining dysprosium, but not the yttrium. He suggested that the most satisfactory methods for freeing holmium from yttrium would be the partial decomposition of the nitrates by heat, or fractional precipitation of the hydroxides with ammonia,2 and by employing the latter method, Holmberg has succeeded in obtaining practi-

cally pure holma. The  $(\beta)$  fractions consist almost entirely of yttrum bromate or ethylsulphate and serve for the isolation of yttrium. For this purpose the fractional decomposition of the nitrates (p 353), the fractional precipitation of the chromates (p. 352), and the fractional precipitation of the basic nitrites (p. 351) are suitable methods, they may be cheaply employed on a large scale, and the first method introduces no volatile reagent. For the final purification Lecoq de Boisbaudian had recourse to fractional precipitation with ammonia, and Cleve to fractional crystallisation of the oxalate from intric acid, R J. Meyer and Wuormen recommend the fractional precipitation of the iodate in nitrie acid solution 4

The (γ) fractions (p. 355) contain yetrium, erbinin, thulium, yeterbium, and lutecum They may be treated by either of the following methods (1.) crystallisation of the bromates, or (ii) crystallisation of the intrates from mitrie acid of density 1.3. The former method is difficult to early out with the ytterbum and lutecum fractions owing to their great solubility, and the temperature should not exceed 15°, but it may nevertheless be prolonged for a sufficient length of time to enable thulium to be isolated in a state of purity 5 The fractions preceding thulium bromate consist of yttrium and erbium bromates, which do not separate appreciably From them erbium

Uıbaın, Compt, rend., 1906, 142, 785; 1909, 149, 37.
 Uıbaın, J. Chim. phys., 1906, 4, 61-2., Ann. Chim. Phys., 1909, [viii], 18, 278.
 Holmberg, Arkir Kein. Min. Geol., 1911, 4, Nos. 2, 10; Zeitsch. anory. Chem., 1911,

<sup>71, 226.

\*\*</sup>Lecoq de Boisbaudian, Compt. rend., 1886, 103, 627; Cleve, ibid, 1882, 95, 1225, G and E. Uthain, ibid, 1901, 132, 136; James and Piatt, J. Amer. Chem. Soc., 1910, 32, 873; Egan and Balke, ibid, 1913, 35, 865, R. J. Meyer and Wuormen, Zeitsch. anory. Chem., 1913, 80, 7.

\*\*James, J. Amer. Chem. Soc., 1910, 32, 517; 1911, 33, 1332.

is best obtained by the fractional decomposition of the nitrates (p. 353), the fractional precipitation of the basic intrites (p. 354), or fractional precipitation with ammonia (p. 352). Pure erbia has not yet been obtained

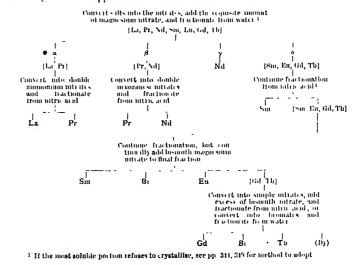
The crystallisation of the simple intrates has been studied by Urbain, it effects the separation of erbinin, thulium, ytterbinin, and Interior slowly, but very satisfactorily, 1 but up to the present only one of these elements has been thus obtained as a pure salt, namely, ytterbium (re neoytterbium) <sup>2</sup> Pure lutecium and celtium salts have yet to be obtained

It remains to be mentioned that small quantities of scandium and thorium may occasionally be present in the more soluble of the yttrium group fractions, the latter element being present owing to its incomplete removal at the commencement. They may be readily eliminated by fractional sublimation of the chlorides, but it is simpler to eliminate the scandium by R. J. Meyer's silicofluoride method, which is described in connection with scandium Any remaining thorium may then be eliminated by the (Chap IX). hydrogen peroxide method (p. 320). It is still simpler to eliminate the scan-

dinm and thorum together as basic thiosulphates (p. 207)

Tabular Summary of General Procedures The following tabular outlines may be of service in helping the reader to follow the preceding descriptions + -

A. Treatment of double sulphates insoluble in sodium or polassium sulphate solution (cerum is supposed to have been removed)



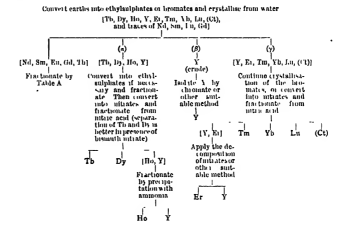
<sup>1</sup> Urbain, J. Chim. phys., 1908, 4, 63, Ann. Chim Phys., 1909, [viii.], 18, 278 Compt. rend., 1907, 145, 759, 1908, 146, 108

2 Blumenfeld and Urbain, Compt. rend., 1911, 159, 323,

3 Urbain, Bourion and Mailland, Compt. rend., 1909, 149, 127.

4 Compute James, J. Amer. Chem. Soc., 1908, 30, 979, 1912, 34, 757.

B. Treatment of crude yttrium earths left in solution from double sulphate treatment-



## HOMOGENEITY OF THE RARE EARTH ELFMENTS.

Cerium.-The homogeneity of cerium was first doubted by Wolf in 1868 1 In 1885 Brauner also concluded that cerum was probably complex, stating that "under certain conditions 'cerium' may consist of a mixture. The nature of this admixture must be ascertained by further experiments "? Ten years later, Brauner stated that further work had substantiated his previous conclusion, and he announced the existence of ordinary cerum (Ce = 140 2), giving a nearly white dioxide, and metacerium, having a higher atomic weight than cerium and forming a dark salmon-coloured oxide. He also mentioned seven methods by which metacerium may be separated more or less completely from cerium 3

In 1895 Schutzenberger announced that cerum was complex, and in 1897, in association with Boudouard, he declared that cerium from cerite was different from the cerium contained in monazite 1. These results, however, were shown to be erroneous by Wyrouboff and Verneurl, who, in the course of their work on the purification of cerra, proved the insufficiency of Schutzenberger and Boudouard's methods for separating ceruum from thorium and the yttrum group, and established the identity of ceria from cente with ceria from monazite.5

In 1900 Drossbach made a prolonged series of crystallisations of ceric

<sup>&</sup>lt;sup>1</sup> Wolf, Amer. J. Sci., 1868, [ii.], 45, 53. Wolf died before his investigations were completed.

Completed.
 Brauner, Trans Chem. Soc., 1885, 47, 879; Monatsh, 1885, 6, 785
 Brauner, Chem. News, 1895, 71, 283.
 Schutzenberger, Cempit rend, 1895, 120, 663, 962; Schutzenberger and Boudouard, 181d, 1897, 124, 181; Boudouard, 181d, 1897, 124, 181; Boudouard, 181d, 1897, 124, 181; Boudouard, 181d, 
ammonium nitrate prepared from 250 kilos, of crude cerous carbonate, but was unable to find any differences among his cerum fractions.1 Since this method was given by Brauner as one by which metacernim could be separated, Drossbach's results throw considerable doubt upon the work of Brauner. The latter chemist returned again in 1903 to the question of the purification of ceria, in an important research on the atomic weight of cerium ence was made, however, to the nature of metacenum 2

At the present time, therefore, it must be admitted that the existence of

metacerium is very problematical

Lanthanum, Praseodymium, Neodymium, and Samarium.—It is generally considered that when the cerum has been removed from the cerium group, four metals remain, namely, lanthanum, praseodymium, neodymium, and samarium. These elements may be quantitatively separated from europium, gadolimum, etc., by the bismuth magnesium nitrate method of Urbain and Lacombe (pp 327, 349)

The homogeneity of each of the elements prascodymnum, neodymnum, and samarium was disputed by a number of chemists very soon after Welsbach's discovery of the complexity of "didynnum" was announced; 3 and in later years others have also denied the elementary character of these substances. Their opinions have been based, however, almost exclusively upon observations of the absorption spectra, and there is little doubt that they are unjustified 5

Demarcay devoted considerable attention to the question of the homogeneity of neodymium and samarium. By fractionating the double ammonium nitrates he isolated a large quantity of the needymnum salt containing only a trace of prascodymnum This material was submitted to a prolonged series of crystallisations. A little praceedynnum and considerable quantities of samarium were thus removed, and more than twenty consecutive ncodymium fractions obtained which presented complete identity from every point of view Demarcay found that needymum thus isolated was the same whether its source was cerite, samaiskite, or mosandrite, and that other methods of fractionation led to the same conclusions. Later he discovered in the fractional crystallisation of the double magnesium intrates a rapid method for the separation of needymum and samarum, and separated several kilograms of the salts of these elements from one another, reducing the intermediate fractions to less than 20 grams. The intermediate fractions showed only the superposed spectra of neodymnum and samarnum, and prolonged fractionation of the samarium salt failed to resolve it into dissimilar parts.

<sup>1</sup> Drossbach, Ber , 1900, 33, 3506.

<sup>&</sup>lt;sup>2</sup> Brauner and Batek, Zeitsch anny Chem , 1903, 34, 103 , Brauner, shid , 1903, 34,

<sup>207.

3</sup> Kruss and Nilson, Ber., 1887, 20, 1676, 2134, 3067; 1888, 21, 585, 2019. Crookes, Chem. News, 1886, 54, 27; C. M. Thompson, ibid., 1887, 55, 227; Bettendoif, Annalen, 1890, 256, 159, 1891, 263, 164; Becqueiel, Ann. Chim. Phys. 1888, [v.i., 14, 170, 257.

4 Denmis and Chamot, J. Amer. Chem. Soc., 1897, 19, 799, Brauner, Proc. Chem. Soc., 1898, 14, 71, Bohm, Die Zerleigharkeit des Pravendyms w. Danst. Sell. End. mit. Hifte einer neuen Trennungsmethode (Halle, 1900); Zeitsch angre. Chem. 1903, 16, 120, Baskerwille and Turentine. J. Amer. Chem. Soc., 1904, 26, 46, Baskerville and Stevenson, ibid., 1904, 26, 51. Although Baskerville and his co-workers speak in a very positive manner of the complexity of prascodymium and neodymium, none of their attempts succeeded in affording any avidence in favour of this view. 

Demarçay's work shows clearly that the ordinary sources of the rare earths contain no significant amount of an element intermediate between needymium and samarium, and argues strongly for the homogeneity of these elements. The subsequent work of various other investigators 1 affords corroborative evidence in favour of Demarcay's conclusions. The homogeneity of prascodymum, however, cannot be so definitely affirmed, though modern work has not yielded any satisfactory evidence for regarding it as complex , 2 moreover, definite information with respect to elements intermediate between prascodymium and neodymium is lacking, although here again modern work points strongly to their non-existence

Lanthana may be defined as the most strongly basic of the cena earths, a colourless oxide that gives rise to salts devoid of absorption spectra. The question of its homogeneity has been studied by Brauner, by Cleve, by Schutzenberger, and by Brauner and Pavlíček. In 1882 Brauner announced the existence of a new earth, intermediate in its equivalent weight and in its basicity between lanthana proper and "didymia." In the same year, Cleve 1 thought he had obtained evidence of the existence of a similar new earth, which he provisionally named oxide of  $Di\beta$ , a few months later, however, he discovered that the spark line  $\lambda 43335$ , characteristic of Di $\beta$ , really belongs to the lanthanum spectrum, the previous measurement of its wave-length by Thalen having been erroneous, and in the following year, in the course of an important research on the atomic weights of lanthanum and "didymum," he definitely declared against the existence of an intermediate element. In 1902 Brauner, in conjunction with Payliček, returned to the subject and maintained the correctness of his earlier work 7. According to Brauner, when lanthana is fractionated by the "basic" method, the least basic fractions contain "a small quantity of an element with a higher atomic weight than that of lanthanum," and the new element "gives to the oxide a very pale buff tint." As other considerations point to the meoriectness of these conclusions, it is desirable that further work on the homogeneity of lanthanum should be undertaken 8

Europium, Gadolinium, Terbium, and Dysprosium.—Europia and gadolima were first isolated in an almost pure state by Demarcay. He was not satisfied, however, that europia was a homogeneous substance, chiefly owing to the fact that it gave rise to salts which exhibited, in his opinion,

See, c q, Muthmann and Weiss, Annalen, 1904, 331, 1. Feit and Prebylla, Zeitsch. anorg. Chem., 1905, 43, 202, 1906, 50, 219, Welsbiel, Sitzingsker K. Akad. Wiss Wien, 1903, 112, 11. A. 1037; Urbain and Lacombe, Compt. rend., 1904, 138, 1166, Urbain, J. Chim. phys., 1906, 4, 119. Eberhald, Zeitsch anorg Chem., 1905, 45, 374. Rech, Zeitsch. wiss Photochem., 1905, 3, 411; Holmberg, Zeitsch anorg Chem., 1907, 53, 83, Baxter and Chapin, J. Amer Chem. Soc., 1911, 33, 1, Baskerville and Stevenson, loc cit.
 Sec, cq., von Scheele, Zeitsch anorg Chem., 1908, 17, 310, Ber., 1899, 32, 409, Welsbach, loc. cit., Lacombe, Bull. Soc. chim., 1904, [in], 31, 570, Chem. News, 1904, 89, 277, Muthmann and Weiss, loc. cit., Feit and Prabylla, loc. cit.; Rech, loc. cit., Brauner, Monatsh., 1882, 3, 486
 Cleve, Compt. rend., 1882, 94, 1528
 Cleve, Lud., 1882, 95, 33.

Cleve, Compt. rend , 1882, 94, 1528

Cleve, thut., 1882, 95, 33.

Cleve, Bull Soc chim., 1883, [ii.], 39, 151, 289

Brauner and Pavlicek, Trans. Chem Soc., 1902, 81, 1243

In 1895 Schutzenberger (Compt. rend., 1895, 120, 1143) declared that by a modification of the method of decomposition of the intrates, lanthana could be resolved into dissimilar parts, the atomic weight of lanthanum varying in the different fractions from 135 to 138.

Brauner and Pavlicek (for. cit.) haveshown, however, that lainthana prepared by Schutzenbarger's method is continuousled with small quantities of vitins. berger's method is contaminated with small quantities of yttila.

abnormally weak absorption spectra 1 The question of the homogeneity of europia and gadolinia was subsequently studied with great care by Urbain, who succeeded, moreover, in isolating terbia and dysprosia and including these earths also within the scope of his researches?

The methods employed by Urbain in effecting the separation of europia, gadolima, terbia, and dysprosia were in nearly all cases crystallisation methods which separate these earths in serial order, ie the order in which they have been mentioned. In each case, the isolation of a pure earth was shown by the constancy of the spark spectrum, absorption spectrum (visible and ultraviolet), and atomic weight throughout a series of fractions, even after further fractionation had been attempted Moreover, the intermediate fractions were reduced to as small amounts as possible in comparison with the quantities of pure earths isolated, and examined for indications of either the presence of other earths or the decomposition of the presumably pure earths obtained. The results were entirely negative.

In the case of europia, Urbam showed that the earth can be freed entirely from samaria and gadolinia, but cannot be resolved into dissimilar prits by prolonged fractionation of the double magnesium intrate in the presence of excess of the corresponding salt of bismuth, that europia isolated by the "bismuth" method from monazite is identical with the earth isolated in the same manner from gadolimite, venotime, or pitchblende, and that europia thus prepared is identical with that isolated by Demarçay's process (p. 311) without the addition of the bismuth salt

In the case of gadolmia, Urbain found that when this earth has been isolated by the "bismuth" method, and shown spectroscopically and by its colour to be free from all but traces of curopia and terbia, further prolonged fractionation of the double nickel nitrates fails to resolve it, but merely accumulates the curopia and terbia in the head and tail fractions respectively, moreover, that gadolima extracted from gadolimte is identical in its properties with gadolinia obtained from xenotinic or pitchblende

The spectral characteristics of terbium were found by Urbain to indicate no signs whatever of the complexity of that element when its separation from gadolimum and dysprosium is effected by any of the following processes: fractional crystallisation of the intrates or double mckel intrates from intric acid and of the ethylsulphates from water or alcohol, fractional precipitation with ammonia, and fractional decomposition of the pitrates by heat - Further, terbia isolated by the ammonia method, after the preliminary application of the ethylsulphate and double mekel intrate processes, is identical with that obtained by crystallising the simple nitrates in the presence of excess of bismuth intrate

Lastly, Urbam found that when dysprosia has been freed as far as possible from impurities by crystallisation of the ethylsulphates, subsequent fractionation of the intrates, while slowly removing the holima and more rapidly eliminating the terbia, fails to indicate that any resolution of the dysprosia \* occurs In fact, forty consecutive dysprosium fractions were obtained by this method of preparation, which were practically identical in all their properties,

A very careful examination of the arc spectra of a number of Urbain's

Demarçay, Compt. rend , 1896, 122, 728 , 1900, 130, 1409 , 1901, 132, 1481 (En);
 ibid., 1900, 131, 343 (Gd).
 See particularly, Urbain, J. Chim phys., 1906, 4, 31, 105, 232, 321; Ann. Chim. Phys., 1909, [viii], 18, 222 , Compt. rend , 1906, 142, 785; 1909, 149, 37

preparations has been made by Eberhard, who concludes that there is no evidence for the complexity of curopium, gadolinium, and terbium, or for the existence of intermediate elements.

From the work of Urbain, then, it would seem legitimate to conclude that europium, gadolinium, terbium, and dysprosium are all well-defined elements. The homogeneity of terbium, however, has been disputed by Auer von Welsbach,3 who, in fractionating the yttria and terbia earths by his double ammonium oxalate method (p 351), obtained indications of its complexity. Welsbach considers torbium to be probably a mixture of thice elements, TbI., TbII., and TbIII. His conclusions, however, must be regarded with considerable suspicion in view of the work of James and Bissel.3 These experimenters found in the fractional crystallisation of the bromates an extremely efficient method for the separation of gadolinium, terbium, and dysprosium, and, on putting this method into practice, isolated only one element from the fractions intermediate between gadolinium and dysprosium. This element exhibited no signs of being complex and answered to the description of terbium as given by Urbain. The balance of evidence is therefore decidedly in favour of the elementary nature of terbium,

The Yttrium Group. - Apart from dysprosium, which has been discussed, the following members of this group receive official recognition in the 1916 table of the International Committee on Atomic Weights holmium, yttrium, erbum, thuhum, ytterbum, and lutecum. The oxistence of Urbain's celtum awaits confirmation.

In the present imperfect state of chemical knowledge concerning this group, it is difficult to form any definite conclusions as to the homogeneity of the above-mentioned elements. By the upholders of the "one band-one element" theory (p. 287), holmium, erbium, and thulium were regarded as complex, and a number of later experimenters have also expressed the opinion that these elements are really composite, basing their conclusions upon a study of the absorption spectra. These conclusions, however, are most probably erroneous, the comparisons of various absorption spectra not having been made under identical conditions 5 In the case of thulium, however, Welsbach 6 has given the wave lengths of the spark lines characteristic of each of the three elements, TmI, TmII, and TmIII, into which he considers thulium can be resolved, and briefly described the three new earths; the homogeneity of thulium is therefore questionable. According to James, the method of fractionation employed by Welsbach is decidedly inferior to the bromate method for the isolation of thulium, being more difficult to operate, involving many more fractions, and effecting the separation more slowly; and yet a prolonged fractionation of thulium bromate, involving 15,000 crystallisations, failed to

Eberhard, Zeitsch. anorg Chem., 1905, 45, 374; Sitzingsber. K Akad Wiss. Berlin, 1906, p. 384, Zeitsch. wiss. Photochem., 1906, 4, 137.
 Auer von Welsbach, Chem. Zeit., 1912, 35, 658.
 James and Bissel, J. Amer. Chem. Soc., 1914, 36, 2060.
 Forsling (Bihang K. Svenska Vet. Akad. Handl., 1902-3, 28, II., No. 1) denies the alementary nature of holmium. Muthmann and Weiss (Annolen, 1904, 331, 1), Welsbach (Monatsh., 1906, 27, 335), and James (J. Amer. Chem. Soc., 1907, 29, 498) consider orbitm to be complex. Marc (Ber., 1902, 35, 2382, Chem. News, 1902, 86, 73) denies the existence of thulium, a conclusion that is unquestionably entoneous.
 Thus Langlet (Aikiv Kem. Min. Gool., 1907, 2, No. 32) contradicts Forsling's conclusions as to the nature of holmium and James (mivrate communication) no longer maintains

clusions as to the nature of holmium, and James (private communication) no longer maintains

that erbium is complex
Welsbach, Monatsh., 1911, 32, 373; Zeitsch anorg. Chem., 1911, 71, 439.

resolve it into dissimilar parts. He accordingly maintains that thulium is really an element 1 Further work upon thulium is therefore necessary.2

It is at present (1916) impossible to state definitely whether there exists any element intermediate between thulum and ytterbunn, but according to Blumenfeld and Urbain 3 this is extremely unlikely.

There is little or no doubt as to the homogeneity of yttium. At one time, however, it was supposed that yttrium is accompanied by another element of atomic weight approximately 97. This supposition was finally disproved by Urbain in 1900, as has been already mentioned, but, while the nature of this supposed element was in doubt, an enterprising firm interested in the manufacture of incandescent gas mantles was enabled to "discover" a new element lucium and to patent methods for the preparation of its salts. The new "element," however, had but a buef existence, its true nature being ascertamed by Crookes'

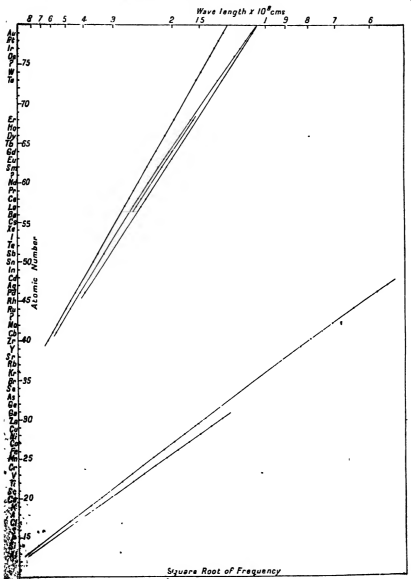
The Cathodic Phosphorescence Spectra and their Bearing on the Homogeneity of the Rare Earth Elements. - The cathodic phosphorescence spectra of the rare carth compounds have been discussed at some length in the preceding chapter (see p 291). From the account there given it will be seen that the study of these spectra clearly indicates that lanthana, prascodymu, neodymua, samaria, europia, gadolimua, terbia, dysprosia, and yttria are substances of the same degree of complexity, and comparable in this respect with alumina, lime, chromium sesquioxide, and other oxides the metallic constituents of which are recognised without question as elementary substances The homogeneity of terbium thus receives additional confirmation. Further, the study of these spectra affords valuable evidence as to the non-existence of any elements between samarium and holmium other than curopium, gadolinium, terbium, and

The X-Ray Spectra and their Bearing on the Homogeneity of the Rare Earth Elements.—The nature of these spectra has already been briefly indicated (see p. 312)

In fig. 37  $^6$  the  $\lambda$  ray spectra of the elements are arranged on horizontal lines spaced at equal distances. The order chosen for the elements is the order of increasing atomic weights, except in the cases of A and K, Ni and Co, I and Te, where this order clashes with that required by the Periodic Further, vacancies have been left, as required by that Classification

classification, between molyhdenum and ruthenum and between tungsten and osmium, and, in addition, one vacancy has been left between needymium and samarium and four vacancies between erbium and tantalium

<sup>1</sup> James, J. Amer Chem Soc., 1911, 33, 1332
2 It may be remarked that in 1909 Urbain stated that he had obtained thulum sufficiently pine for its principal characteristics to be determined with accuracy, although the fractions still contained a little erbain or ytterbain according to their position. No mention was made of any indications of its complexity. Urbain's method of preparation, namely, fractional crystallization of the initiates, differs from the methods of Welshach and James (see Urbain, Ann Chem Phus, 1909, [viii], 18, 279).
3 Blumenfeld and Urbain, Compt. rend., 1911, 159, 323.
4 For further information on this point, and references to the literature, see p. 355.
5 See Barrière, Chem. Nics., 1896, 74, 159, 212. Crookes, thid., 1896, 74, 259, R. Fresenius, ibid., 1896, 74, 269, Shapleigh, ibid., 1897, 76, 41
The "elements" Losmium and neclosmium (Kosmann, Zeitsch Elektrochem., 1896-7, p. 279. cf. Winkler, Br., 1897, 30, 13) were of a similar nature to lucium.



F10. 37.—High-frequency spectra of the elements (Moscley).

arrangement of the elements is equivalent to assigning to successive elements a series of successive characteristic integers. The integer N for Al, the thirteenth element, has been taken as thirteen. The corresponding integers for the other elements are indicated in the diagram and are spoken of as their atomic numbers. As abscisse, the square roots of the frequencies (r) of the lines have been plotted.

It is clear from the diagram that when the values of N are plotted against those of vi, the points he on a series of smooth curves which approximate closely to straight lines. Such a remarkable relationship between the X-ray spectra of the elements cannot be regarded as accidental, it is undoubtedly of great theoretical importance. The deductions that may be drawn from it in connection with the rare earths are as follows: (i) an element, presumably intermediate in atomic weight between Nd (1413) and Sm (1504), has yet to be discovered, (ii) terbinim and holimum must be regarded as elementary substances, and (iii) four rare earth elements come between erbinim (1677) and tantalum (1815). With regard to (iii), it follows that thulium cannot be a mixture of three elements, as Welsbach maintains, since ytterbinim and lutecium must account for two of the available places, if, on the other hand, thulium is really elementary, the existence of celtium becomes very probable, the four blanks being presumably filled in order by thulium, ytterbium, lutecium, and celtium

Granting, then, the existence of celtium, it would appear that all the raio carth elements at present recognised are really elementary substances, and that only one other remains to be discovered. That this unknown element should come between needynnum and samarium is, however, surprising in view of the negative result obtained by Demargay in his search for such an element (p. 359). It may be that the missing element is extremely rare, and occurs in some rare earth immerals but not in others, as celtium appears to do, with this possibility in view, search is now being made for it by James.<sup>2</sup>

Firsther work on the X-ray spectra of the rare earth elements will be awarted with considerable interest

### DETECTION AND ESTIMATION OF THE RARE EARTHS 3

Qualitative Analysis. --The analytical reactions of the rare earth elements are summarised in the accompanying table, in which the reactions of scandium, titanium, zircommin, and thorium have been included for purposes of comparison. A number of special tests for cerium are known, but characteristic reactions for the other rare earth elements are lacking; the formation of a blue adsorption compound of basic lauthanum accetate and rodine, for instance, which has been proposed as a test for lauthanum, is not

stances ordinarily recognised as elements

2 Prof. James, private communication to the author. It may be mentioned here that m working up the rare earths from certain minerals, Prof. James has observed that small quantities of a radioactive substance concentrate in the similium fractions. Further work on this point is in progress.

<sup>1</sup> That is to say, they have as much right to the title of "clement" as have the sub-

on this point is in progress.

For further information on this subject, see R. J. Meyer and C. Hauser, Die Analyse der seltenen Erden und der Erdsauren (Enke, Stuttgart, 1912), Mellor, A Treatise on Quantitalize Analysis (C. Griffin & Co., Ltd., 1913). On the analysis of cerium alloys see also Arnold, Zeitsch. anal. Chem., 1914, 53, 496, 678; Rolling, Ann. Chim. Applicata, 1914, 2, 366.

# THACTIONS OF THE RARE EARTH ELEMENTS COMPARED WITH THOSE OF SCANDIUM, ZIRCONIUM, AND THORIUM.

|          | •                                                                             |                                                                                        |                                                            |                                                                                                |                                                                                                         |                                               |                                                                                                       |                                                           | _            |
|----------|-------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|------------------------------------------------------------|------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------|-----------------------------------------------|-------------------------------------------------------------------------------------------------------|-----------------------------------------------------------|--------------|
|          | Reagent                                                                       | Cerum Group,<br>including Cerii                                                        | Quadrivalent<br>Cerium Ce <sup>17</sup>                    | Yttrum Group                                                                                   | Scandium                                                                                                | Titanium                                      | Zuconum                                                                                               | Тһогит                                                    |              |
| i        | ım, so-                                                                       | Hydrovide pre-<br>cipitated 2                                                          | Hydrovide pre-                                             | Hydroxide pre-                                                                                 | Hedrovide pre-                                                                                          | Hydrovide pre-                                | Hydroxide pre-<br>cipitated                                                                           | Hydrovide j<br>cipitated                                  | pre-         |
|          | nyurokide<br>Nodlum, potass-<br>num, or ammonium<br>carbonate                 | Carbonate pre-<br>cipitated, slightly<br>soluble in excess of                          | Excess of reagent<br>produces a clear,<br>solution         |                                                                                                | Same as strium                                                                                          | Hydrovide pre-                                | Hydroxide pre-<br>cipitated, soluble<br>in ammonium car-                                              | Basic salt pre-<br>cipitated, soluble<br>in excess of re- | ple<br>re-   |
|          | Precipitated lead                                                             | reagent 3<br>No precipitate                                                            | Hydroxide pre-                                             | excess of reagent. No precipitate                                                              | •                                                                                                       | Hydroxide pre-                                | Donate<br>Hydroxide pre-<br>ciritated                                                                 | agent<br>Hydroxide 1<br>civitated                         | pre.         |
| -        | carlionate<br>Sodium, potass-<br>lum, or ammonium                             | Double sulphates<br>precipitated                                                       | Precipitate only                                           | No precipitate                                                                                 | Precipitate only                                                                                        | No precipitate                                | Precipitate ouly                                                                                      | Precipitate only<br>with K salt                           | ylu,         |
| 9, 9     | sulphate (excess of saturated solution) Solum acetate (horing solution) Solum | No precipitate                                                                         | Hasic acetate pre-<br>cipitated<br>Hydrovide pre-          | No precipitate No precipitate                                                                  |                                                                                                         | Hydroxide pre-<br>cipitated<br>Hydroxide pre- | Hydrovide pre-<br>cipitated                                                                           | Basic acetate<br>precipitated<br>Hydroxide pre-           | tate<br>pre- |
| į-       | sulphate (boiling<br>solution)<br>Hydrogen per-<br>oxide in intitral or       | No precipitate                                                                         | cipitated Reduced to Celli                                 | No precipitate                                                                                 | pnace precipitate                                                                                       | Xillowish - red                               | Hydrated per-<br>oxide precipitated 4                                                                 |                                                           | per-         |
| 00       | ammoniumovalate                                                               | Ovalate precipitated, slightly selluble in mineral acids or excess of ammonium ovalate | Reduced to Celli,                                          | Oxalate precipi-<br>tated, rather more<br>soluble in acids or<br>amnonium oxalate<br>than Ceii | Oxilate precipitated, sparingly soluble in acids, readily in aminimitation mention mention oxilate.     | No precipitate                                | Basu oxalate pre-<br>cipitated, readily<br>soluble in ovalic<br>acid, ammonium<br>oxalate, or mineral | Same as scan-                                             | , a          |
| 6        | Tartaric acid                                                                 | No precipitate,<br>and precipitation<br>with amnionia in-                              | Reduced to Cell.                                           | No precipitate, int with aumonia precipitates double                                           | Sanie as yttrium                                                                                        | Same as Celli                                 | acids<br>Same as Cem                                                                                  | Same as Celli                                             |              |
| 2        | Hydrofluoric acid                                                             | hibited Precipi- Fluoridible in alkali flucrides Appreciably solibile in hot nimeral   | :<br>:                                                     | tarrates<br>same as Cem                                                                        | Fluoride precipitated, soluble in alkalı fluorides Nuch less soluble in mineral acuds than the Cuit and | No precipitate                                | No precipitate                                                                                        | Same as scan-<br> drum                                    | can-         |
| <b>=</b> | Iodic acid                                                                    | Iodate precipi-<br>tated, renuily sol                                                  | . Isolate precipi<br>tated, spaningly<br>soluble in intric | Same as Cem                                                                                    | Y salts Iodate precipi tated, furly casily soluble in nitric                                            | Incomplete hydrolysis                         | Indate precipatively practically insoluble in nitro                                                   | Same as                                                   | L            |
|          |                                                                               |                                                                                        | 1-4-1                                                      | 0                                                                                              | 9 Comme he decembe conduces to come hadrounds on exposure to sir (n. 342)                               | yana to corio hadana                          | de ou exposure to air                                                                                 | (n 38%)                                                   |              |

1 See also L Haber, Monath, 1897, 18, 687

2 Cerous hydroxide oxidises in air, forming an alkali perceric carbonate (p. 401)

3 The cerous solution oxidises in air, forming an alkali perceric carbonate (p. 401)

4 From zeronyl chloride solution, but not from nitrate or sulphate solutions unites ammonia is also added.

a specific reaction of that element, for it is given by praseodymium and possibly by other rare earth elements (see pp. 410 and 416)  $^1$ 

Special reactions of cerium salts are as follows:

(i.) To a slightly acid or neutral solution of a cerous salt, hydrogen peroxide and an excess of ammonium acetate are added, and the inixture warmed to 40°-60°. A yellowish brown precipitate of basic percene acetate is produced even when only I part of cerium is present in 10,000 of solution (Hartley's reaction).2

(11) Very dilute ammonia is added, drop by drop, to a solution of a corous salt until a permanent precipitate just commences to form. A few drops of hydrogen peroxide are added and the mixture well shaken. A yellow precipitate or coloration is produced even when only traces of ceria

are present and other rare earths present in quantity.3

(iii.) A solution of a cerous salt is warmed with very concentrated potassium carbonate solution, and a little hydrogen perovide added to the clear solution, the solution turns yellow even when only a trace of cerum

(iv) A solution of a cerous salt is boiled with an ammoniaeal solution of When traces ammonium taitrate, it turns dark brown owing to oxidation of cerum are sought, the oxidation may be accomplished by means of hydrogen peroxide 5

(vi) The lead peroxide, sodium bismuthate, and ammonium persulphate methods mentioned later in connection with quantitative analysis (p. 373)

may be utilised as qualitative tests for ceruin

(vii.) Ammonated silver intrate is added to a neutral solution of a cerous salt and the mixture warmed, a brown coloration or black precipitate is produced according to the amount of cerum present.6

(vm ) The cerum solution is made alkaline with sodium hydroxide, evaporated to diviness, and a drop of a solution of strychimic in excess of sulphuric and added. The solution turns blue or violet in colour and then

becomes red 7

The usual methods for separating the rare earth elements from other metals in the course of analysis utilise the insolubility of (i) the hydroxides in ammonia, (ii ) the fluorides in dilute hydrofluoric acid, and (iii ) the ovalates m dilute mineral acids. The following process may be employed. The metals that form insoluble chlorides and sulphides insoluble in dilute hydrochloric acid are removed, hydrogen sulphide expelled from the filtrate, and ammonum chloride and ammonia then added. The precipitate is dissolved in hydrochloric acid, the solution evaporated to dryness, a decided

<sup>1</sup> Praseodymnum gives a yellowish given borar bead in the ovidising, and a given bead in the reducing flame, in odymnum gives a violet bead in the reducing flame. The colours are seen only with cold beads (Milbauer, Zeitsch anal Chem., 1907, 46, 657). On the marochemical tests for the lare earths, see Poirt-Escot and Conquet, Compt. rend., 1900, 130, 1316, Behrens, Arch. Nierland., 1901, [11], 6, 67, Rec. trav. chim., 1905, 23, 413; R. J. Meyer, Zeitsch. anny Chem., 1902, 33, 113

4 Haitley, Trans. Chem. Soc., 1682, 41, 202. Dennis and Magee, J. Amer. Chem. Soc., 1894, 10, 619.

5 Leccul de Boisbandian. Compt. rend., 1885, 100, 605. Cleve. Bull. Soc. chim., 1885.

<sup>1894, 16, 619.</sup>Lecaq de Botsbaudian, Compt read, 1885, 100, 605, Cleve, Bull Soc cham, 1889
[1] 43, 57, Dennis and Magee, loc. ett., Marc. Ber., 1992, 35, 2370
Job, Compt rend, 1898, 126, 246, R. J. Meyer, Zedisch anory Chem, 1904, 41, 94.
Wirth, Chem Zett., 1913, 37, 773.
W. Biltz and Zimmermain, Ber., 1907, 40, 4979.
Sonnenschein, Ber., 1870, 3, 631, Bull. Soc. cham, 1870, [ii.], 14, 201, Plugge, Arch. Pharm., 1891, 229, 558; Dennis and Magee, loc. cat.

11

excess of hydrofluoric acid added to the residue, and the liquid boiled for a minute or two. The residue of fluorides is filtered off, heated with concentrated sulphuric acid, the sulphates dissolved in water, and any barium, strontium, or calcium sulphate removed by filtration The rare earth elements are then precipitated from the filtrate by the addition of ammonia. They are free from titamium and zircomium, but contain any scandium and thorum originally present 1

For the qualitative separation of the rare carths themselves, so far as this is usually attempted, the methods given for their quantitative separation may be followed

Quantitative Analysis.—In the absence of other elements, a rare earth clement is usually estimated by precipitation as oxalate, followed by ignition to the oxide Precipitation is preferably made from a boiling, dilute intrate solution, slightly acidified with nitric acid, by the addition of oxalic acid, 2 the oxide finally obtained from a chloride solution is very hable to contain traces of basic chloride, presumably derived from oxalochloride co-precipitated with the oxalate 1. The oxalate precipitate should be allowed to stand for several hours (preferably overnight) before it is filtered. The oxide finally obtained should be strongly ignited, ceria, for instance, as obtained from cerous oxalate by heating over a Bunsen burner, loses 0 5-0 7 per cent, of its weight when ignited over the blowpipe 4. In all but three cases, the oxide obtained is the sesquioxide, M<sub>2</sub>O<sub>4</sub>, ceriuin, however, leaves the dioxide, CeO<sub>2</sub>, and praseodymium and terbium are obtained as peroxides which may be assumed 5 to have the composition M107, as alternatives, their perovidic oxygen may be determined indunctrically, or the peroxides may be reduced to sesquioxides by heating in hydrogen. In weighing the sesquioxides it should be borne in mind that they take up moisture and carbon dioxide from the air fairly rapidly

The use of an alkah oxalate as procipitant instead of oxalic acid has been carefully studied by Baxter, Griffin, and Daudt,6 who have found that, in neutral solution, co-precipitation of alkali oxalate occurs to a very appreciable extent, except with sodium oxalate, which is only carried down in small amounts. The contamination increases with rise of temperature, but may usually be greatly diminished if precipitation is conducted in the presence of a quantity of a strong acid considerably more than equivalent to the oxalate added Contamination with ammonium oxalate is, of course, ununportant for the gravimetric determination of the rare earth, but is as serious as contamination with any other oxalate when it is desired to carry out the estimation volumetrically by titrating the rare earth oxalate.

A convenient alternative to the precipitation as oxalate is the precipitation as sebacate, effected by adding ammonium sebacate to a boiling, neutral solution of the nitrate 8 Precipitation as hydroxide is also satisfactory when

See A. A. Noyes, Bray, and Spean, J. Amer. Chem. Soc., 1908, 30, 481.
 See, e.g., Lenhen, J. Amer. Chem. Soc., 1908, 30, 577
 See, e.g., Job, Compt. rend., 1898, 126, 246; Matignon, Ann. Chem. Phys., 1906, [vin.], 8, 215.
 Wytonbolf and Verneurl, Ann. Chem. Phys., 1906, [vin.], 9, 297
 Welsbach, Monatsh., 1885, 6, 477, Baxter and Griffin, J. Amer. Chem. Soc., 1906, 28, 1684; Urbain and Janitech, Compt. rend., 1908, 146, 127.
 Baxter and Griffin, J. Amer. Chem. Soc., 1906, 28, 1681, Baxter and Daudt, ibid., 1908, 30, 563
 For the titiation, see the references cited on p. 242.
 Whittemore and Janies. J. Amer. Chem. Soc., 1912, 34, 772, 1913, 35, 129.

Whittemore and James, J Amer. Chem. Soc., 1912, 34, 772, 1913, 35, 129.

precipitation is made by means of ammonia from a nitrate solution precipitation of basic chloride when the chloride is used leads to results that are slightly high, on the other hand, adsorption of alkali leads to very high results if sodium or potassium hydroxide is used as precipitant? Matignon has recommended the determination of the rare earths by weighing them as the anhydrous sulphates 2

Separation of the Rave Earth Elements from other Metals - It might be anticipated that the separation of the rare earth elements from sodium or potassium could readily be accomplished by precipitation with ammonia or oxalic acid Such, however, is not the case, for appreciable quantities of alkali compounds are co precipitated A satisfactory separation is afforded however, by precipitation with ammonium schaeate from a boiling, neutral solution of the intrate.3 The fluoride method also appears to be satisfactory A quantitative separation from lithing may be effected with oxalic acid if the precipitation is made in a cold solution in the presence of ammonium chloride.

The first steps taken in effecting the quantitative separation of the raic earth elements are usually the same as those followed in qualitative work. metals which form chlorides or sulphides insoluble in dilute hydrochloric acid are removed in the usual manner, the filtrate freed from hydrogen sulplinde, and the rare earth elements then precipitated as hydroxides by the addition of ammonium chloride and ammonia hon, aluminium, chromium, glicinium, zirconium, thorium, mannin, and titanium may be co-precipitated, while an almost complete separation from zinc, menganese, nickel, cobalt, magnesnim, and the alkali and alkaline earth metals is obtained. The separation may be rendered more efficient, if necessary, by solution of the precipitate in acid and reprecipitation with ammonia — In rare earth immeral analysis, chromium, zinc, mckel, cobalt, barnin, strontinin, and the alkali metals are seldom encountered, while titamim is usually removed at the commencement

The rare earth elements are separated from the other metals contained in the ammonia precipitate by means of oxalic acid When the cerum group predominates, the following conditions are most sintable. The precipitate is dissolved in hydrochloric acid, the concentration of the acid being adjusted so as to be between 0.25N and 0.5N, and 60 e.e. of acid being used for a gram of mixed earths. Precipitation is then effected by the addition of sufficient oxalic acid for the final hquid to contain 3 grams of oxalic acid per 100 c.c., and the whole maintained at about 60' for twelve hours. The best conditions when the yttimm group predominates require to be investigated

The care earth elements are thus separated as ovalates, but still in association with thorium. With respect to the separation from the other metals, it has been observed that the rare earth ovalates are appreciably soluble in solutions of uranyl salts, and also, though to a less extent, in ferric and aluminum salts, complex manyloxilates, etc. being produced, hence a considerable excess of oxalic acid has to be added to effect a quantita-

<sup>&</sup>lt;sup>1</sup> T. O. Smith and James, J. Amer. Chem. Soc., 1914, 36, 9c6., Chem. News, 1914, 1. O. Smith and James, J. Amer. Chem. Soc., 1914, 36, 900. Chem. News, 1914, 29, 219
 Matignon, Ann. Chem. Phys., 1906, [vm.], 8, 247
 Winttemore and James, J. Amer. Chem. Soc., 1912, 34, 772 (VA., 1913, 35, 129 (La., Nd). In the case of Y.—K. mixtures, double prorphation is necessity.
 Winttemore and James. Jul. 1919, 24, 779 (VA., 279)

Mittenore and James, while, 1912, 34, 772 (Y).

Hauser and Witth, Zeitsch anal Chem., 1908, 47, 389.

Hauser, Zeitsch anal. Chem., 1908, 47, 677.

Dittrich, Ber., 1908, 41, 4373.

tive precipitation, and the determination of small amounts of rare earths in uranium minerals is rendered very difficult.1

The separation of the rare earth elements from thorum is most conveniently dealt with in a separate section, but at this point the methods for opening up rare earth minerals for analysis may be mentioned. As a matter of fact they have already been enumerated at the beginning of this chapter (see p. 318). Silicates may, if necessary, be fused with sodium carbonate, but as a rule they may be completely decomposed by evaporation with aqua regia. Phosphates, such as monarite and xenotime, may also be decomposed by fusion with sodium carbonate or fusion mixture, and removal of the alkali phosphate produced by extracting with dilute sodium hydroxide, although monazite is usually treated with sulphuric acid or sodium bisulphate. The columbotantalates and related innierals may be attacked by one of three methods. (i.) The mineral is fused with an excess of sodium bisulphate. The mass is extracted with water, the solution largely diluted, and boiled for several hours in a flask fitted with a reflux condenser. If titanium can then be detected in the liquid, the free acid present is partially neutralised with ammonia and the boiling continued until the titanium is entirely precipitated. The liquid is then filtered, it contains all the rare earth elements, thorium, zinconium (part), uranium, etc., while the residue may contain columbic, tantalic, titame, tingstie, stannie, and silicie acids, zircomum (part), iron (part), and sparingly soluble sulphates or basic sulphates such as those of lead, mercury, bismuth, etc. (11) The finely powdered mineral is decomposed by means of hydrofluoric acid (see p. 319) (m) The inneral, contained in a porcelain or silica boat, is placed in a combustion tube and heated to dull redness in a stream of the vapour of disulphur dichloride Columbium, tantalum, titanium, and tungsten are thus removed as volatile chlorides or oxychlorides, and collected in dilute nitic acid, together with some or all of the volatile chlorides of iron and aluminum. When decomposition is complete, the residual chlorides in the boat are cooled in a stream of hydrogen chloride and their analysis effected in the manner already described 5

Traces of rare earths are occasionally found in silicate rocks, and may be determined as follows. The powdered rock is partially evaporated with hydrofluoric acid several times, the insoluble fluorides or silicofluorides separated by filtration and converted into sulphates by means of concentrated sulphuric acid The filtered solution of the sulphates in dilute hydrochloric acid is precipitated with ainmonia, the precipitate dissolved in hydrochloric acid, and the rare earths precipitated as oxalates

According to Whittenore and James (J. Amer. Chem. Soc., 1912, 34, 772), yttimum is satisfactorily separated from from aluminum, and magnesium by precipitation with oxalic sold in a cold solution to which summonium chloride has been added. Smith and James (15td.) 1913, 35. 563) have shown that precipitation with oxalic acid in a slightly acid solution gives a satisfactory separation of neodyimum from titanium, manium, glueinum, zirconium, and barum.

and baruun.

<sup>2</sup> Sec, e.g., Chesneau, Compt rend. 1911, 153, 429.

<sup>3</sup> Sec, e.g., Glaser, J. Amer. Chem. Nov., 1896, 18, 782. Chem. Zeit., 1896, 20, 612;

Fresenus and Hintz. Zeitsch andel. Chem., 1896, 35, 525. Hintz and Weber, thid, 1897, 36, 27, 676; Benz. Zeitsch angew. Chem., 1902, 15, 297; Johnstone, J. Soc. Chem. Ind., 1914, 33, 55. For another method of attacking phosphates, see Bailot and Chauvenet, Compt. rend., 1913, 157, 1153

<sup>4</sup> On the separation from tungsten, see also Wunder and Schapus. Ann. Chim. anal., 1913, 18, 257

<sup>5</sup> Hicks, J. Amer. Chem. Soc., 1911, 33, 1492

<sup>8</sup> W. F. Hillebrand, The Analysis of Silicate and Carbonate Rocks (\*\*Bull. U.S. Geol. Survey." 1910, No. 422), p. 143, where an alternative procedure will also be found.

Survey," 1910, No. 422), p. 143, where an alternative procedure will also be found.

Separation of Thorium from the Rare Earth Elements. - The mixed oxalates of the rare earth elements and thorum are converted into the intrates by heating on the steam-bath with a mixture of ordinary concentrated intric acid and the fuming acid 1. The liquid is evaporated to dryness and the residue dissolved in 100 c c of 10 per cent ammonium intrate The solution is warmed to 60'-80' C, and the thornin precipitated by the addition of pure 3 per cent hydrogen perovide. The precipitate is filtered, washed with hot dilute ammonium intrate, ignited wet in a platinum crucible, and weighed as thoria, ThO, This method gives very good results, for great accuracy, the moist precipitate may be dissolved in intricacid and the thorium reprecipitated as peroxide.3

Alternative methods are the thiosulphate, iodate, pyrophosphate, hypophosphate, fumarate, meta introbenzoate, lead carbonate and sebacate pro cesses mentioned earlier in this chapter (see p. 321, where references to the literature will be found), precipitation as basic thorum acetate,4 precipitation of the thornm as hydroxide by certain weak organic bases, volatilisation of the thornum as the chloride, etc - The thiosulphate method is the oldest and one of the most convenient. It is rather tedious to carry out when results of a high degree of accuracy are required,7 but for many purposes the procedure may be simplified. Precipitation should be effected from a dilute solution of the mixed chlorides after having evaporated it to dryness or added ammonia to expel or neutralise free acids, and the sodium thiosulphate added should be at least equivalent to the total rare earths present. After boiling for fifteen minutes only a trace of thorann remains in solution, but, unless the thora constitutes some 90 per cent or more of the total earths, the precipitate contains small amounts of earths other than thoria, which must be removed by a second precipitation  $\Delta$  third precipitation is seldom necessary

Separation of the Rare Earth Elements from One Another.-A general solution of this problem is not yet known. As a rule the cerium is determined and an approximate separation of the remaining elements into the cerum and yttrium groups effected. Two methods of procedure are available (i) the cerum is separated first by a suitable graymetric method. and (it) the separation into the ceruin and yttimm groups is carried out first, and the cerum subsequently estimated either gravimetrically or

volumetrically 5

to the presence of prascodymum droxide in the oxides weighed.

The error caused by the prascodymum may be channeled by heating the oxides to redness in hydrogen, but only when certa has been previously removed, for certa loses in

weight when heated in hydrogen.

<sup>1 (</sup>If Borelli, Gazzella, 1909, 39, 1–125

2 According to R. C. Wells of Non-them Ind., 1910, 29, 1301), this results from the balancing of slight positive and negative errors

3 Wytorboil and Venneul, Bull Societim, 1897, [m], 17, 679; 1898, 19, 219; Complexed 1898, 126, 340, 127, 412, Ann-Chem Ibys., 1905, [vin], 6, 489. Benz, Zedsch. angew Chem., 1902, 15, 297.

4 Mingaye, Records God Survey N.S. Wales, 1909, 8, 276.

5 Mins Jelleison, J. Amer. Chem Nov., 1902, 24, 540, Hartwell, ibid., 1903, 25, 1128.

6 Bourion, Ann Chim. Phys., 1910, [vin], 21, 109

7 For the necessary procedure, see White, Lature on Thorum and its Compounds (The Institute of Chemistry, 1912); cf. Johnstone, J. See Chem Ind., 1911, 33, 56, and the references cited on p. 321.

references cited on p. 321.

<sup>\*</sup> When procedure (ii.) is employed and the cerium determined volumetrically, the volumetric analysis is made after the weight of the total rare earths of the cerium group, including the ceria, has been ascertained—the ceria earths other than ceria may then be calculated. This latter result, however, will be slightly in excess of the true value owing

Separation of the Cerium.—Assuming the thorium to have been removed by the peroxide method, the filtrate is treated in a dish with excess of ammonia and hydrogen peroxide, the liquid boiled, and then the whole is evaporated to dryness on the steam-bath. The residue is carefully heated until ammonium intrate commences to decompose, or else dried at 110°; the perceric hydroxide is thus converted into ceric hydroxide. The residue is dissolved in strong nitric acid and the cerium separated as basic ceric autrate-sulphate as described on p 333, about 0.1 gram of ammonium sulphate being used to 0.5 gram of mixed rare earths. The precipitate is filtered off and washed with hot dilute ammonium intrate. To the filtrate 0.5 gram of sodium acetate and about 0.5 grain of ammonium persulphate are added, and the liquid boiled. The remaining cerium is thus precipitated, together with traces of the other elements. The two precipitates are ignited together at a white heat, and the cerum weighed as ceria, CeO, 1 In mineral analysis the result is then calculated to cerous oxide, Ce<sub>2</sub>O<sub>3</sub>, since the cerium immerals are cerous compounds

In addition to the preceding process the bromate method described on p. 334, and Mosander's chlorine method, outlined on p. 336, may be employed <sup>2</sup>
Separation of the Cereum and Ytterum Groups — This separation is usually effected by the double sulphate method. The rare carth elements, having been freed from thorium, are suitably treated to obtain a very concentrated and nearly neutral solution of the chlorides. A large excess of a saturated solution of potassium sulphate is added, together with a few grams of the powdered salt, and the whole allowed to stand for twelve hours, with occasional agitation. The crystalline precipitate is washed with saturated potassium sulphate solution, reconverted into the chloride, and the separation repeated From the combined filtrates the yttma earths are precipitated as hydroxides, and then converted successively into intrates, oxalates, and oxides. The total ceria earths are extracted from the precipitated double sulphates in a similar manner. The weights of the mixed oxides give approximately the amount of ceria and yttria earths present 3 Sodium sulphate is not so satisfactory a precipitant as the potassium salt

An alternative method has been proposed by James and Smith 4 solution of the mixed intrates is first prepared. This is evaporated to drynoss, treated with 25 c.c. of concentrated nitric acid saturated with bismuth magnesium intrate, and then a further 10 grains of bismuth magnesium intrate and I gram of magnesium intrate added. The whole is warmed and stirred until most of the salt has dissolved, it is then set aside for twenty-four hours. The precipitate is collected in a Gooch crucible and washed with intric and saturated with bismuth magnesium intrate

In this method the metals of the yttrium group remain in solution, while those of the cerum group are precipitated almost quantitatively 5

<sup>1</sup> Wyrouboff and Verneuil, Compt. rend., 1899, 128, 1331 . Ann Chem. 1 hys , 1906,

<sup>[</sup>vin.], 9, 345.

For details of the former, see James and Pratt, J. Amer. Chem. Sec., 1911, 33, 1326; for details of the latter, see Johnstone, J. Soc. Chem. Ind., 1914, 33, 55., Hauser and Writh, Zeitsch. and Chem., 1909, 48, 679; Browning and Roberts, Amer. J. Sci., 1910, [iv.],

<sup>23, 15.
39, 15.
39, 16.
39, 16.
4</sup> James and T. O Smith, J Amer. Chem. Sα., 1912, 34, 1171.
5 Precisely how samarium, earoprum, gadolimum, terbium, dysprosium, and holmium behave is not known.

Volumetric Estimation of Cerium - Several volumetric methods are available whereby cerium may be accurately determined in the presence of the other rare earth elements. They are all based upon the conversion of cerons into ceric salts or rice rersa.

(1) The mixed intrates, in cold, concentrated intric acid solution, are treated with an excess of lead peroxide and shaken initil oxidation of the cerous salt is complete. In the filtered solution the ceric salt is titrated with standard hydrogen peroxide until the colour of the cene salt just

$$2\text{Ce}(NO_3)_1 + H_2O_3 = 2\text{Ce}(NO_3)_3 + 2HNO_3 + O_3$$

(ii ) The lead peroxide of method (i ) is replaced by bismuth tetroxide, oxidation being effected in cold intrie acid (I vol. of acid to I of water). The solution is diluted, tiltered through asbestos, excess of standard hydrogen peroxide added, and the excess titrated with standard potassium permanganate 2

(iii ) The oxidising agent used is sodium bismuthate, and oxidation effected in boiling, dilute sulphuric acid (1 of acid to 1 of water) in the presence of a gram or two of ammonium sulphate. The filtered solution is reduced with standard ferrous sulphate, excess of which is titrated with potassium permanganate 4

(iv) Oxidation to the cerie state is effected in boiling, dilute sulphuric acid by means of aumonium persulphate, the titration is then effected in the cold liquid by the use of hydrogen perovide and potassium permanganate.1

This method is largely employed. It gives good results, but requires careful attention to points of detail

(v.) The oxidation is effected by means of potassium ferricyanide in the presence of alkalı hydroxide -

$$Ce_2(SO_1)_3 + 2K_3Fe(CN)_6 + 8KOH = 2Ce(OH)_4 + 2K_4Fe(CN)_6 + 3K_2SO_4$$

The ceric hydroxide is removed by filtration and the potassium terrocyamide in the filtrate titrated with polassium permanganate in the presence of sulphune acid  $^{\circ}$ 

(vi) The neutral solution, containing the cernini as cerous salt, is added, with constant shaking, to a definite volume of hot, standard potassium permanganate containing excess of magnesia in suspension, until the permanganate is decolorised — 6

$$\begin{aligned} &3\text{Ce}(\text{NO}_{4})_{4} + \text{KMnO}_{4} + 4\text{MgO} + 8\text{H}_{2}\text{O} \\ &= 3\text{Ce}(\text{OH})_{4} + \text{MnO}_{2} \\ &2\text{H}_{2}\text{O} + 4\text{Mg}(\text{NO}_{4})_{2} + \text{KNO}_{3}, \end{aligned}$$

Job, Compt. rend., 1899, 128, 101, Ann. Chem. Phys., 1900, [vii.], 20, 205,
 Chesneau, Compt. rend., 1911, 153, 129. of Gibbs, Amer. J. Ser., 1-61 [ii.], 37, 354;
 Buhrig, J. prakt. Chem., 1875, [ii.] 12, 231
 Waegner and Muller, Ber., 1903, 36, 282, 1732, of Gibbs, Amer. Chem. J., 1893,

<sup>15, 546.

3</sup> Metzger, J. Amer. Chem. Soc., 1909, 31, 523, Metzger and Hendelberger, ibid, 1910,

<sup>Metzger, J. Amer. Curm. Soc., 1808, 32, 822, 823, 842
Von Knoire, Zeitsch. angen. Chem., 1897, 10, 685, 717. Ecr., 1900, 33, 1924,
Power and Shedden, J. Soc. Chem. Ind., 1990, 19, 696
Strohlen h (Mat. Techn. Versuchsamtes, 1912, 1, 45. J. Soc. Chem. Ind., 1913, 32, 199) oxidises with chloric acid
Browning and Palmer, Amer. J. Net., 1908, [v. ], 26, 83, Chem. News, 1908, 98, 106;
Zeitsch anurg Chem., 1908, 59, 71
R. J. Meyer and Schweitzer, Zeitsch. anorg. Chem., 3907, 54, 104, cf. Brauner, Chem. News, 1895, 71, 283, Lenher and Meloche, J. Amer. Chem. Soc., 1916, 38, 66.</sup> 

(vii.) When ceria is digested in an atmosphere of carbon dioxide with a warm solution of potassium iodide in concentrated hydrochloric acid, it is quantitatively reduced to the cerous state and the liberated iodine may be titrated with sodium thiosulphate -- 1

$$2\mathrm{GeO}_2 + 2\mathrm{KI} + 8\mathrm{HCl} = 2\mathrm{GeCl}_3 + 2\mathrm{KCl} + 4\mathrm{H}_2\mathrm{O} + \mathrm{I}_2.$$

This reaction, however, is of no practical value for the estimation of ceria in mixtures of earths.2

(viii) Small quantities of ceria present in thoria may be determined colorimetrically.3

Quantitative Analysis by Spectroscopic Methods —(1.) Spark Spectra.—When the relative degrees of persistency of the lines in the spark spectrum of, say, lanthanum chloride solution, are known, it is clear that the results may be utilised to advantage for the purposes of quantitative analysis. A dilute chloride solution of the substance containing lanthanum is prepared, of known concentration, and its spark spectrum examined. A first approximation to the lanthanum content of the solution being thus obtained, closer and closer approximations to the true value may then be obtained in successive trial experiments. The feasibility of such a method of analysis was demonstrated as long ago as 1884 by Hartley, 1 at the present time, though, the necessary data for the spark spectra of the rare earth elements are almost enturely lacking 5

(n.) Absorption Spectra. - Two general methods may here be employed for the estimation of those rare earth elements that give rise to absorption spectra.

In the first method, the absorption produced by the element in the substance to be analysed is recorded photographically and compared with a series of photographs of the absorption produced by the element at a series of known concentrations and under the same conditions of experiment

In the second method, the amount of absorption exerted by the substance is measured quantitatively at those regions of the spectrum where its absorption is most characteristic. The necessary photometric measurements are effected by the use of a spectrophotometer, for a description of which the reader must be referred elsewhere The method of calculation may, however, be briefly outlined.

Suppose that a layer of absorbing medium is used of thickness d, the layer being bounded in part by two parallel faces, on to one of which homogeneous light 6 of intensity Io falls normally. If the light emerges from the opposite face with intensity I, then

$$I/I_0 = a^d$$

where a is a constant (the absorption coefficient) the value of which depends upon the nature of the medium and the wave-length of the light. If d' is

<sup>&</sup>lt;sup>1</sup> Bunsen, Annalen, 1853, 86, 265, 1858, 105, 49; Browning, Ame. J. Sci., 1899, [iv.], 8, 451, Chem. News, 1900, 81, 30, 41, Zet/-ch. anory. Chem., 1899, 22, 297. Brauner, "Std., 1903, 34, 207

<sup>2</sup> Marc, Ber., 1902, 35, 2370; R. J. Meyer and Koss, Ber., 1902, 35, 3740, Mengel, Zet/sch. anory. Chem., 1899, 19, 71.

<sup>3</sup> Ben., Zet/sch. anory. Chem., 1899, 19, 71.

<sup>4</sup> Hartley, Ph.l. Trans, 1884, 175, 49, 325.

<sup>5</sup> For La, Ce, and Y, see Polloft and Leonard, Sci. Proc. Roy. Dubl. Soc., 1908, 11, 257, and the table of wave lengths given on p. 307.

<sup>6</sup> Le. light of one wave-length.

the particular value of d for which  $I/I_0=1/10$ , then 1/d' is called the "extinction coefficient" and denoted by a. It follows that  $a=10^{-a}$  and

$$-\log_{10}(1/I_0) = ad$$

In accordance with Beer's Law, if c', c'', c'''... be the concentrations of the absorbing substance, and the corresponding thicknesses of absorbing layer for the light intensity to fall from  $l_0$  to l be d', d', d''', then c'd' = c''d'' = c'''d'' = . If, then, the extinction coefficients of the solutions be a', a'', a''', ..., it follows that

$$c'/a' = c''/a'' = c'''/a''' = ... = \Lambda,$$

or, in general terms,

$$c = aA$$

where  $\Lambda$ , called the "absorption ratio," is a constant that may be determined once for all for any particular absorbing substance in a given solvent

Suppose, now, that the concentration (c) of neodymnum in an aqueous solution of rare earth chlorides is required the extinction coefficient (a) is measured by the spectrophotometer at a particular part of the spectrum where the absorption ratio (A) for aqueous neodymnum chloride is known. From the relationship  $c=a\Lambda$  the required concentration can then be mimediately calculated

It has been already pointed out in the preceding chapter that the absorption exerted by a rare earth salt is modified by the presence of a salt of another rare earth element, even though the second salt gives no absorption spectrum. Hence measurements of concentrations effected by observations upon absorption spectra only give approximate results when mixtures are examined, i.e. in the only cases where one need resort to the process at all <sup>1</sup>

<sup>&</sup>lt;sup>1</sup> For further information, see Bahr and Bunsen, Annalm, 1866, 137, 30. Muthmann and Statzel, Rev. 1899, 32, 265;. G. and H. Kruss, Koloremetre and quantilative Spektrolandlys (Voss, Leipzig 2nd ed., 1909), Thorpe, A Internary of Applied Chemistry (Longmans, 1912, 1913), vol. v. article "Spectrum Analysis", Meyer and Hauser, Die Analyse der seltenen Erden und der Erdsauren (Enke, Stuttgart, 1912).

# CHAPTER XII.

# THE RARE EARTH ELEMENTS (continued).

## THE CERIUM GROUP.

The rare earth elements included in this group are five in number, namely, lanthanum, cerum, praesodymium, neodymium, and samarium. The chemistry of these elements is fairly well known, the commercial treatment of monazite sand for thorium having enabled large quantities of the earths of the cerum group to be placed at the disposal of scientific investigators.

The isolation of ceria from the mixed ceria earths is a comparatively simple operation. The separation of the remaining members of the group is most conveniently effected by the fractional crystallisation of the double ammonium, magnesium, or manganese intrates, as described in the preceding chapter.

The points of difference between the cerium and yttrium groups have been already stated (see p. 219)

In the following account of the ceruin elements and their compounds cerium itself is dealt with first, the other elements being then taken in their serial order.

# CERIUM.

Symbol, Ce. Atomic weight, 140 25 (0 = 16).

Cerium is the most abundant of the rare earth elements. Its occurrence, listory, atomic weight, homogeneity, and analytical chemistry have already been discussed in Chapters X, and XI.

The separation of certa from the other rare earths and its purification have been discussed in some detail in Chapter X. (see p. 332). On a commercial scale pure cernum salts are obtained by (i) the permanganate method (p. 335), (ii.) the basic nitrate method (p. 332), or (ii) the certa ammonium nitrate method (p. 336), the procedures already outlined being suitably modified for technical work. Only a small proportion of the cernum available (in monazite residues) is worked up into pure cernum salts.

Metallic cerum is obtained by the electrolytic methods previously described (p. 230).

Terium is a metal which resembles steel in appearance and takes a high polish. Its density is 6 92 at 25°. It is malleable and highly ductile,

<sup>&</sup>lt;sup>1</sup> The properties of cerium as here given are substantially those stated by Hirsch, J. Ind. Eng. Cham., 1911, 3, 880, 1912, 4, 955, Trans. Amer. Electrachem Soc., 1911, 29, 57. See also Muthmann and Kraft, Annalen, 1902, 325, 261, Muthmann and Weiss, Annalen, 1904, 331, 1; Hillebrand and Norton, Poyg. Annalen, 1875, 155, 633; 156, 466

may be readily cut with a knife, and can be machined fairly well, although there is a tendency to buckle. Its ultimate strength is 9 kilos per square mm. The metal is paramagnetic, its magnetic susceptibility at 18° being 15  $\pm 10^{-6}\,\mathrm{c\,g\,s}$  units per gram  $^{1}$ . Cerum is a fairly good conductor of heat, but a poor conductor of electricity, its resistance being 71 6 micro ohins per cm cube at the ordinary temperature. The following values have been obtained for the specific heat -2

| Mendeléeff | 9        | spec | ht= 0 05 | atomic ht | 7 01  |
|------------|----------|------|----------|-----------|-------|
| Hillebrand | 0°- 100° | ٠,,  | =0.0118  | ,,        | -6.28 |
| Hirsch .   | 20°-100° |      | -0.05112 | **        | -717  |

Hirsch's value was obtained by using the differential steam calorimeter and operating with 70 grams of cerum. The atomic heat of cerum thus appears to be unusually high. Cerum melts at 635° (Husch), 623 (Muthmann and Weiss).

Cerum takes from air at 160°, burning with even greater brilliancy than magnesium and evolving much heat. A shower of sparks is produced by striking the metal with a flint. When a lump of cerium is kept warm in a closed bottle, a black powder slowly forms on the surface, and when the bottle is opened the powder inflames spontaneously. Cermin burns brilliantly m chlorine at  $210^\circ\text{--}215$ , and in bromine at  $215^\circ$ – $220^\circ$ , it also combines directly with hydrogen, introgen, iodine, sulphur, selemini, tellinium, arseine, antimony, etc. It reduces carbon monoxide and carbon dioxide with the separation of carbon

Cerum preserves its lustre in dry, but tarmshes in moist an - It is very slightly attacked by cold water, but in boiling water a slow evolution of hydrogen occurs. At the ordinary temperature ethyl and amyl alcohols, chloroform, earbon tetrachloride, and concentrated solutions of sulphuric acid, sodnim and ammonium hydroxides have no action on cermin, dilute hydrogen peroxide, ammonium and potassium chloride slowly attack it, and the action of hydrochloric or nitric acid, both dilute and concentrated, and dilute sulphuric acid is moderately vigorous. At boiling temperatures the immeral acids, dilute or concentrated, attack the metal rapidly, except concentrated sulphuric acid.

According to Huseh, cerum is best purified by heating it with boiling mercury in a long non pipe. The cerium amalgamates with the mercury, and the impurities may be skimmed off from the molten amalgain amalgam is placed in a magnesia vessel, the whole placed miside a larger quartz vessel, and the amalgam heated very strongly in vacuo to drive off the mercury

Alloys.—Cerum alloys readily with a large number of metals. Many of the alloys thus obtained are hard and buttle, differing therefore from pure cerum, which is soft and malleable. Accordingly, small particles are easily detached from the alloys by friction, and when the alloys are rich in cerium the heat produced is more than sufficient to ruse the temperature of the particles above 160°, the ignition-point of cerium. Hence numerous alloys rich in cerium are pyrophorie. Some of them, eg non-cerium alloys, are of

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1911, 14, 637.

Mendelécs, Bull load Set Petrograd, 1870, p 445, Hillebrand, Pogy Annalen, 1876, 158, 71, Ilusch, loc. cit.

<sup>1</sup> M. Owen, Ann Physik, 1912, [iv ], 37, 657, Proc K 1kad Wetensch Amsterdam,

commercial importance owing to this property, being utilised in the manufacture of automatic lighters, etc. The alloy of cerum, lanthanum, etc., known as "Mischinetall," is a valuable reducing agent (see p. 230).

Sodium alloys with cerium quietly, forming a hard and somewhat pyrophoric alloy that oxidises on exposure to air Silver and gold alloy readily with cerium; the silver alloys are hard and brittle, but the gold alloys are fairly soft.<sup>2</sup> The copper alloys have been systematically examined and four compounds discovered, viz Cu<sub>0</sub>Ce (m p 935°), Cu<sub>1</sub>Ce, Cu<sub>2</sub>Ce (m p 820°), and

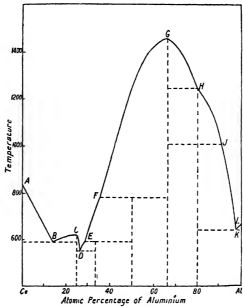


Fig. 38.—The system communication administration of the system communication of the system of the sy

CuCe. Between 55 and 85 per cent. Ce the alloys are pyrophoric, and with more than 30 per cent. Ce they soon disintegrate

Calcium and cerium alloys are white and very hard, they are stable in air, do not disintegrate, and emit brilliant sparks when struck. They may be prepared from the metals, or by reducing cerous fluoride with calcium.4 Magnesium and cerium alloys are white in colour and extremely brittle.

<sup>1</sup> See Kellermann, Die Ceritmetalle und ihre pyrophoren Legierungen (Knapp, Halla 

The alloys rich in cerium form excellent flashlight powders. Those containing 60-75 per cent. of cerium may be finely powdered, and form excellent reducing agents, eg for preparing vanadium from its oxide. Alloys containing more than 75 per cent, of cerum are difficult to powder without catching fire. The alloys have been systematically examined by Vogel, who has described the compounds Ce<sub>4</sub>Mg (m p 632°), CeMg (m p 738°), CeMg<sub>3</sub> (m p 780°), and CcMg<sub>0</sub> The compound Cc<sub>4</sub>Mg breaks up into Cc and CcMg at 497', the compounds CcMg and CcMg, form solid solutions with one another. Zinc alloys with cerium almost explosively. The alloy is hard, brittle, and pyrophone, but ordises in the air \*\*Cachinum\* behaves similarly to zine \*\* Mercury readily alloys with cernin \*\* The amalgams, which are liquid at ordinary temperatures when only 1 or 2 per cent of cernin is present, decompose water with the evolution of hydrogen and oxidise quickly in the air, spontaneous combustion readily occurring when 8-10 per cent of cerium is present 2

Aluminium and cerum alloys have been studied by the thermal and microscopic methods. The equilibrium diagram, due to Vogel, is shown in fig. 38 3 No solid solutions are formed. Five compounds exist, two only being stable at the melting point. Ce, \(\frac{1}{4}\) (mp. 614), Ce, \(\frac{1}{4}\), CeAl, CeAl, (eAl, CeAl, The compound CeAl, exists in two forms, the transition-point being 1005. The form stable at ordinary temperatures is probably thombic. Its density is \(\frac{1}{193}\). The cerum almost meaning are stable to an analysis are stable to an analysis of the cerum almost and the content of the cerum almost are stable to an analysis of the cerum almost are stable to an analysis of the cerum almost are stable to an analysis of the cerum almost are stable to an analysis of the cerum almost are stable to an analysis of the cerum almost are stable to an analysis of the cerum almost are stable to an analysis of the cerum almost are stable at ordinary temperature. stable to air and water unless the cerum is more than 75 per cent. The maximum hardness is 6, corresponding to CeAl. This compound is very resistant towards acids, but burns brilliantly

For other cerum alloys, see the subsequent volumes of this series Thermochemistry of Cerium. — The following results have been

```
\begin{bmatrix} Ce \end{bmatrix} + \begin{pmatrix} O_2 \end{pmatrix}\begin{bmatrix} Ce \end{bmatrix} + \begin{bmatrix} A1 \end{bmatrix}
  [CeO<sub>2</sub>] . . . . . [CeAI<sub>1</sub>] . . . .
   + 224.6 Cals
  + 121 1
     [Ce(OH)<sub>i</sub>] + H<sub>2</sub>SO<sub>i</sub>aq
   \mathrm{Ce}(\mathrm{SO}_1)_2\mathrm{aq}
   Ce<sub>3</sub>(SO<sub>4</sub>) aq + H O<sub>4</sub>q + (O<sub>5</sub>) + 29 95 ,,
Ce<sub>3</sub>(SO<sub>4</sub>) aq + H SO<sub>4</sub>qq + (O<sub>2</sub>) + 33 58 ,,
\begin{array}{l} 2 \big[ \mathrm{Ce}(\mathrm{OH})_3^\top \mathrm{O} \cdot \mathrm{OH} \big] + 3 \mathrm{H}_2 \mathrm{SO}_{\mathrm{p}} \mathrm{u}_1 \\ 2 \mathrm{Ce}(\mathrm{SO}_1)_3 \mathrm{u}_1 + \mathrm{H}_2 \mathrm{O}_2 \mathrm{u}_1 \\ - \big[ \mathrm{Ce}(\mathrm{OH})_3^\top \mathrm{O} \cdot \mathrm{OH} \big] \end{array} -
   + 20.39
  = \left[ \left( \left( OH \right)_{1} \right) \right] + \left( O \right) .
```

# CEROUS COMPOUNDS

The cerous salts are derived from the basic oxide Ce<sub>2</sub>O, and, if derived from colourless acids, are themselves colourless. Their aqueous solutions are devoid of absorption spectra.

In chemical properties the Gerous salts resemble the salts of lanthanum very closely, except for the fact that they may be oxidised to cene salts. The equivalent conductivities, \(\lambda\), of cerous salts are in harmony with the view that

<sup>1</sup> Vogel, Zeitsch anneg Chem., 1915-91, 277

2 Muthmann and Beck, Anneha., 1901., 331, 46, Husch, Ic., cit.

3 R. Vogel, Zeitsch anneg. Chem., 1912., 75, 41.

4 Muthmann and Beck, lor eit

5 Muthmann and Wess., Innalen., 1901-331, 1 (Gr.O.), Muthmann and Beck, ibid.,

1904., 331, 46 (Co.Al.); Pissaijewsky, J. Russ Phys. Chem. Soc., 1900., 32, 609. The data given for ceric and perceric hydroxides refer to the hydroxides as ordinarily precipitated.

they are derived from a fairly strong triacid base. The following data hold for a temperature of 25° (v = dilution in litres per grain-equivalent). — 1

Numerous cerous salts have been described by John 2 and others The methods for the conversion of cerous into cene compounds and vice versa are discussed later (p 391).

## CERIUM AND HYDROGEN.

Cerous hydride, Cell, has been described (see p. 251). The combination of cernum and hydrogen proceeds readily even at 310°, but is best effected at 450°, at which temperature the dissociation pressure of the hydride does not exceed 1 mm 3 The hydride is spontaneously inflammable.4

# CERIUM AND THE FLUORINE GROUP.

Cerous fluoride, CeF, is said by Brauner to be obtained by carefully heating ceric fluoride to dull redness 5. The hemilydrate, 2CeF, H,O, is obtained by double decomposition of Cerous fluoride forms no compound with potassium fluoride.7

Cerous chloride, CeCls.—The methods of preparation and properties of the anhydrous salt are described on p 252. Bourion's sulphur chloride method, starting with ceric oxide, is very convenient The chloride is a white, crystalline, hygroscopic solid, readily soluble in water

Anhydrous cerous chloride readily absorbs ammonia. The following compounds have been prepared, and their dissociation pressures measured .- 8

An aqueous solution of cerous chloride may be prepared by heating ceric hydroxide with concentrated hydrochloric acid, chlorine being evolved. On saturating the cold solution with hydrogen chloride, orthorhombic crystals (a:b:c=0 8083.1.1.4419) of the heptahydrate, CeCl<sub>3</sub> 7H<sub>2</sub>O, separate. A triclinic salt (a:b:c=1.1580:1:0.8635; a=91°3′,  $\beta$ =114° 3′,  $\gamma$ =88° 12′) has also been described, isomorphous with lanthanum chloride heptahydrate, but it is said to be the hexahydrate, CeCl<sub>3</sub>.6H<sub>2</sub>O.<sup>10</sup>

Aufrecht, Inaugural Dissertation (Beilin, 1904), cf. Holmberg, Arkiv Kem. Min.

<sup>1</sup> Aufiecht, Inaugural Dissertation (Beilin, 1904), cf. Holinberg, Arker Rem. Sala.

Geol., 1903, r, 1.

2 Jolin, Bull. Soc. chim., 1874, [n ], 21, 533.

3 Zhukov, J. Russ. Phys. Chem. Soc., 1913, 45, 2073.

4 Dafeit and Miklauz, Monatsh., 1912, 33, 911.

5 Brauner, Ber., 1831, 11, 1914; Trans. Chem. Soc., 1882, 41, 68. See also p. 393.

6 Jolin, Bull. Soc. chim., 1874, [ii.], 21, 533.

7 Puchin and Baskov, J. Russ. Phys. Chem. Soc., 1913, 45, 82.

8 Barre, Compt. rend., 1913, 156, 1017.

9 Definis and Magee, J. Amer. Chem. Soc., 1804, 16, 619. Zeitsch. anory. Chem., 1894, 7, 250; cf. Jolin, Bull. Soc. chim., 1874, [n ], 21, 533, Lange, J. prakt. Chem., 1861, 82, 129; R. J. Meyer and Koss, Ber., 1902, 35, 3740.

10 Fock, Zeitsch. Kryst. Min., 1894, 22, 32.

The following double salts are known, in addition to those containing metals of Groups IV. and V. and organic bases -

```
\begin{array}{l} {\rm CeCl_3~4HgCl_3~10H_2O~,^1~CeCl_3.3Hg(CN)_2.8H_2O~^2~CeCl_3.AuCl_3~13H_2O.^{1.4}~\\ {\rm CeCl_3~2PtCl_2~10H_2O~;^7~CeCl_3~PtCl_1~12H_2O.^{1.4~5.5}~\\ \end{array}
```

Cerous oxychloride, CeOCI (\*) has been described by several chemists \* Cerous bromide, CeBr, -The anhydrous salt, a white, hygroscopic mass, has been prepared.9 A hydrate and an auribromide, CeBi, AuBi, 8H2O (John), are known

Cerous iodide, Cel, -The hydrate, Cel, 911,0, forms colourless crystals

which turn brown in the air 10

Cerous perchlorate, Ce(ClO<sub>1</sub>), 8H<sub>2</sub>O or 9H<sub>2</sub>O (1), is a deliquescent, crystalline salt, which may be prepared from cerous sulphate and barium perchlorate 11

Cerous bromate, Ce(BrO<sub>3</sub>), 9H<sub>2</sub>O, must be crystallised from aqueous solution by concentrating in vacuo at 35° and then cooling. It melts at 49°, and decomposes at slightly higher temperatures - Further, it slightly decomposes in aqueous solution 12

Cerous iodate, Ce(10,1), 211,0, is readily obtained by double decomposition as a white, amorphous, sparingly soluble precipitate 13 At 25°, one litre of saturated solution contains 1 61 grams of the anhydrous salt.11

# CERIUM AND THE OXYGEN GROUP.

Cerium sesquioxide or cerous oxide, Ce2O p so readily oxidises to form cerum dioxide that cerous salts of volatile oxyacids yield ceria when ignited in the air. The sesquioxide can be prepared by the reduction of the dioxale with hydrogen at 150 atmos pressure and at a temperature of about 2000° (' (ct. p. 393) 15. In small quantities and mixed with much cern and a little carbon, it is produced by the ignition of cerous oxidate in vacuo or in hydrogen. It is also produced in poor yield by heating cern

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1 John, loc cit
       2 Alen, Orfices Sienska Vet Akad Forhandl., 1876, 33, Nos 8, 9, Bull Soc. chim.,
1876, [u.], 27, 365
      76. [n. ], 27, 305

3 Holzmann, J. prakt. Chem., 1861, 84, 76

4 Holzmann, Jahresber., 1862, p. 1.25

5 Martynac, Ann. Chem. Phys., 1873, [n. ], 30, 67

6 Cleve, Bull. Sw. chim., 1874, [n. ], 21, 196

7 Nilson, Ber., 1876, 9, 1056. See also p. 254

8 Wohler, Annalen, 1867, 144, 251, Enk. Juhresber., 1870, p. 319, Dulier Compt. rend., 855, 107, 882.
1885, 101, 882.

See p. 255 , also Robinson, Proc. Roy. Soc., 1884, 37-150
Lange, J. prakt. Chem., 1861, 82, 129
John, loc. cit., Morgan and Cahen, Trans. Chem. Soc., 1907, 91, 175
See p. 256, and Rammelsberg, Popp. Annalen. 1842, 52, 84
Rammelsberg, Popp. Annalen., 1838, 44, 557, Holzmann, J. prakt. Chem., 1858, 75, 15
John, Loc. cit.
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Rammelsheig, Forg. Innaten, 1838, 44, 551, 1102mini, 3 pract view. 1838, 75, 321, John, loc. ct.
Rimbach and Schubert, Zeitsch. physikal. Chem., 1909, 67, 183
Kowbery and Pring, Proc. Roy. Soc., 1916, A, 92, 276. According to Guertlei (Zeitsch. amory, Chem., 1904, 40, 247). Cerium dovide is quantitatively reduced to the sesquioxide when heated to a white heat in the reducing atmosphere of a petroleum injector-furnace.
Mosandei, Phil. May., 1843. [in.], 23, 241; Delafontanic, Arch. Ser. phys. nat., 1865, 22, 38; Bulnig, J. prakt. Chem., 1876; [in.], 12, 209. Popp. Annaten., 1864, 131, 759; Steiba, Ann. Chim. Phys., 1904, [vin.], 2, 209. See also Beringei, Annaten, 1842, 42, 138; Rammelsbeig, Poyy. Annaten, 1859, 105, 40; Winkler, Ber., 1891, 24, 878.

with zinc at the boiling-point of the metal, but it cannot be separated from the other products of the reaction.\(^1\) The simplest method yet known for its preparation consists in heating ceria with metallic calcium and dissolving out the lime and excess of calcium in aqueous ammonium chloride at - 10° C. sesquioxide is thus obtained as a yellowish-green powder which readily absorbs oxygen even at ordinary temperatures and burns easily when gently heated.2

Cerous hydroxide, Ce(OH), is obtained as a white precipitate by the addition of ammonia or alkali hydroxide to a solution of a cerous salt, or by the action of water on cerum carbide, CeC2. It may be dried without losing its white colour if air is completely excluded.3 It cannot, however, be further dehydrated to cerous oxide, for the latter, as it is produced, attacks the water

present and oxidises.

Cerous hydroxide, like the other hydroxides of the rare earth elements, is a strong base. It readily oxidises in the air, becoming first violet (p 393) and finally yellow, owing to its conversion into cenic hydroxide. Oxidising agents effect the change more rapidly (p. 396). Owing to the ease with which it is oxidised, cerous hydroxide is a strong reducing agent, reducing cupric salts to cuprous, moreuric salts to mercurous, etc.4

Cerous sulphide, Co<sub>2</sub>S<sub>3</sub>, may be prepared by heating ceric oxide to bright redness in a stream of dry hydrogen sulphide, the oxide, contained in a porcelain boat inserted in a porcelain tube, being heated until its weight ceases to alter (see p 259). An alternative method is to use cerous sulphate instead of ceric oxide and heat to 750°-800° in hydrogen sulphide.<sup>5</sup>

Cerous sulphide is a dark reddish-purple solid of density 5.02 at 11° (Muthmann and Stutzel). When heated in the air it roasts to cerous sulphate and ceric oxide, but in hydrogen or nitrogen it is stable till 1100°-

1500°, when it melts and decomposes

Cerium persulphide, Co,S, is prepared by heating cerous sulphate to 580°-600° in a stream of dry hydrogen sulphide, the preparation being followed by weighing from time to time. The persulphide is a dark yellowish-brown, crystalline powder, moderately stable in air and cold water. It dissolves in cold hydrochloric acid, an odour of hydrogen persulphide being first noticed and subsequently a precipitation of sulphur. The compound therefore appears to be a polysulphide. At 720° it decomposes into the sesquisulphide and sulphur; it may be reduced to the sesquisulphide by heating in hydrogen at 400°.6

Cerous oxysulphide, Ce<sub>2</sub>O<sub>2</sub>S, may be prepared by heating cerium dioxide to redness in a current of wet hydrogen sulphide. It is a goldenyellow solid which has not been obtained quite free from impurities.7

Cerous sulphite, Ce<sub>2</sub>(SO<sub>3</sub>)<sub>3</sub> —Hydrates of this salt with 3H<sub>2</sub>O and 9H<sub>2</sub>O

are known.8

Cerous sulphate, Ce (So4) .- The preparation and general properties of the anhydrous sulphate are given on p. 260. In 100 parts of ice-cold water

Burger, Ber., 1907, 40, 1652.
 Dennis and Mager, J. Amer. Chem. Soc., 1894, 16, 649.
 Dannica, Compt. rend., 1913,

<sup>157, 214.</sup> W. Biltz and Zummermann, Ber , 1907, 40, 4979 , Bailneil, Att. R. Accad. Lincei, 1907, [v.], 16, 1, 395, ...

See p. 259; also W. Biltz, Ber, 1908, 41, 3341.

W. Biltz, loc. ct.

Steiba, Ann. Chim. Phys., 1904, [viii]. z, 193.
 See p. 260; and Batčk, Zeitsch anorg. Chem., 1905, 47, 87.

40 parts of cerous sulphate can be dissolved. The specific heat (0° to 100°) of the anhydrous sulphate is 0·1168  $^2$ 

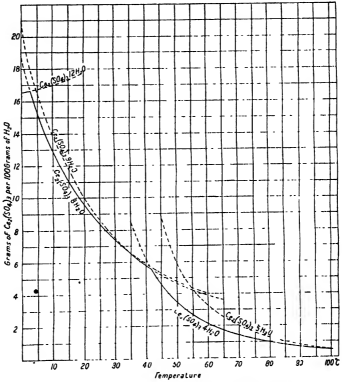


Fig. 39 -Solubility of the hydrates of cerous sulphate in water

Cerous sulphate forms numerous hydrates. The existence of hydrates with III<sub>2</sub>O, 5H<sub>2</sub>O, 8H<sub>2</sub>O, 9H<sub>2</sub>O, and 12H<sub>2</sub>O is definitely established, but the existence of the hydrate with 61120 is very doubtful 3

<sup>1</sup> Muthmann and Rolig. Zeitsch. anory Chem., 1898, 16, 450. On the densities of aqueous solutions of cerous sulphate, see Biauner, Trans. Chem. Soc., 1888, 53, 257.

2 Nilson and Petterson, Ber., 1880, 13, 1459, Compt. rend, 1880, 91, 232.

3 The hydrates of cerous sulphate have been studied in detail by Koppel (Zeitsch anorg. Chem., 1904, 41, 877), who gives a critical review of callier work on the subject. The other references are as follows. Other, Pong. Annalen, 1837, 40, 404; Manignac, Mem. Sci., phys. nat., 1855, 14, 201; Inn. Chim. Phys., 1873, [iv.], 30, 56, Androwicz, J. prakt., Chem., 1880, 80, 16; Hermann, wid., 1813, 30, 184; 1864, 92, 113; Wolf, Amer. J. Sci., Chem., 1880, 11; J. 45, 53; Wing, wid., 1870, [ii.], 46, 356; John, Bull. Soc. chem., 1844, [ii.], 21, 533; Muthmann and Rolig, loc. cit., Wyrouboff, Bull. Soc. chem., 1901, [iii.], 25, 105; Bull. Soc. franc. Min., 1901, 24, 105; Kraus, Zeilsch. Kryst. Min., 1901, 34, 307.

The various hydrates may be obtained as follows .- The dodecahydrate, Co2(SO1)8 12H2O, separates from a concentrated solution of cerous sulphate over sulphuric acid at 0° C. in the form of tiny needles. The enneahydrate, over supported activate of the form of tiny neededs. The emeasurates,  $Ce_2(SQ_4)_3.9H_2O$ , is prepared by evaporating the aqueous solution at  $40^\circ-45^\circ$ , and forms hexagonal pusms. The octahydrate,  $Ce_2(SQ_4)_3.8H_2O$ , is best prepared by warning to  $45^\circ$ - $50^\circ$  a solution of the sulphate that was saturated at  $0^\circ$  C. The pentahydrate,  $Ce_2(SQ_4)_3.5H_2O$ , is obtained by preparing a cold saturated solution and heating it to  $70^\circ-100^\circ$ , and forms monoclinic prisms. The tetrahydrate,  $Ce_3(SQ_4)_3.4H_2O$ , is conveniently prepared by agitating the appealydrate with a little water at  $70^\circ$  for 48 hours and dying the mass enneahydrate with a little water at 70° for 48 hours, and drying the mass at the same temperature For densities, see p. 260.

The crystals of the penta- and enneahydrates have been already described (p. 261). The octahydrate has been generally regarded as crystallising in the orthorhombic system, but according to Wyrouboff, whose view is adopted by Groth, the crystals are trichine and devoid of all elements of symmetry  $(a \cdot b \cdot c = 1.0650 \cdot 1 : 1.1144, a = 90^{\circ} 52', \beta = 90^{\circ} 40', \gamma = 91^{\circ} 45')$ . Rammelsberg has described a monochnic octahydrate, isomorphous with the octahydrates of

The solubilities of the various hydrates, in grams of anhydrous sulphate per 100 grams of water, are given in the accompanying table (Koppel) and represented graphically in fig 39, from which the stable hydrate at any temperature is readily seen It will be noticed that solubilities have been determined even

where the hydrates are metastable in contact with the solution -

| t° C.<br>12H₀O | 0°<br>16 56 | 18 8<br>17 52 | 19·2°<br>17·70 |             |             |                |             |                |
|----------------|-------------|---------------|----------------|-------------|-------------|----------------|-------------|----------------|
| t° C.<br>9H₀O  | 0°<br>20 98 | 15°<br>11 87  | 30°<br>7 35    | 45°<br>5 13 | 50°<br>4 67 | 60°<br>3·88    | 65°<br>3 60 | 0              |
| t° C.<br>8H₀O  | 0°<br>19 09 | 15°<br>11 06  | 20 5°<br>9 53  | 30°         | 40°<br>5 95 | 50°<br>4 79    | 60°         | v              |
| t° C.<br>5H₂O  | 45°<br>8.83 | 60°<br>3 25   | 70°<br>1·93    | 80°<br>1·21 | 90<br>0 81  | 100 5°<br>0:17 |             |                |
| t° C           | 35°<br>8:50 | 40°<br>6.04   | 50'            | 57°<br>2 34 | 65"<br>1.88 | 70°<br>138     | 82°<br>1:01 | 100 5°<br>0·43 |

The acid sulphate, Ce(IISO<sub>4</sub>)<sub>3</sub>, is known (see p. 263).

The following double salts with alkali sulphates have been described -

 $\begin{array}{l} \text{Ce}_2(\text{SO}_4)_3, \text{K}_2\text{SO}_4, 2\text{H}_2\text{O}, ^6\text{2Ce}_2(\text{SO}_4)_3, 3\text{K}_2\text{SO}_4, ^2 \cdot \text{and with } 8\text{H}_2\text{O}, ^6\text{Ce}_2(\text{SO}_4)_3. \\ 2\text{K}_2\text{SO}_4, ^2\text{H}_2\text{O}, ^4\text{Ce}_2(\text{SO}_4)_3, 3\text{K}_2\text{NO}_4, ^2 \cdot ^4\text{Ce}_2(\text{SO}_4)_3, 5\text{K}_2\text{SO}_4, ^6\text{Ce}_2(\text{SO}_4)_8, \text{Na}_2\text{NO}_4, 2\text{H}_2\text{O}, ^{8,6}\text{Ce}_2(\text{SO}_4)_8, (\text{NH}_4)_2\text{SO}_4, 8\text{H}_2\text{O}, ^{3,4,6}\text{ and anhydrous}; ^{4,7}\text{Ce}_2(\text{SO}_4)_3, 5(\text{NH}_4)_2\text{SO}_4, ^6\text{Ce}_2(\text{SO}_4)_3, (\text{NH}_4)_2, (\text{NH}_$ 

<sup>1</sup> For crystal measurements, see Maugnac, loc. cit., Wytouboff, loc. cit., Kraus, loc. cit.; Desclouseaux, Mim. l'Indidut, 1858, p. 111; Viba, Zeitsch. anory Chem., 1904, 39, 283; Groth, Chemische Krystallograpie (Leipzig, 1906-10), vol. n., Rammelsberg, cited by

Groth, op cit.

Bermann, loc cit.

Czudnowicz, loc. cit.

John, loc. cit.

Beringer, Annalen, 1842, 42, 138.
 Barie, Compt. rend., 1910, 152, 871.
 Wolff, Zeitsch. anorg. Chem., 1905, 45; 89.

They are sparingly soluble in water and practically insoluble in saturated solutions of the corresponding alkali sulphates.

The diminution in the solubility of cerous sulphate in water, caused by the presence of various alkali sulphates, is as follows (data represent grams of anhydrous salt present in solution per 100 grams of water):--1

| Temp                                         | 16° C.                                | Temp                                         | . 19° C.                             | Temp.                                              | 16° ('.                                       |
|----------------------------------------------|---------------------------------------|----------------------------------------------|--------------------------------------|----------------------------------------------------|-----------------------------------------------|
| K <sub>2</sub> SO <sub>4</sub>               | ('e <sub>4</sub> (SO <sub>4</sub> ),  | Na <sub>2</sub> SO <sub>4</sub> .            | ('e/(SO <sub>4</sub> ),              | (NII <sub>4</sub> ) <sub>2</sub> SO <sub>4</sub> , | ( 'e <sub>2</sub> ( SO <sub>4</sub> ),        |
| 0 00<br>0 18<br>0 51<br>0 73<br>1 29<br>2 50 | 10 75<br>0 96<br>0 13<br>0 25<br>0 04 | 0 00<br>0 33<br>1 09<br>1 39<br>1 70<br>3 59 | 9.65<br>0.64<br>0.09<br>0.06<br>0.03 | 0 00<br>3 46<br>9 32<br>19 24<br>15 62<br>72 81    | 10 75<br>1 03<br>0·78<br>0 75<br>0 50<br>0·04 |

Cerous ammonium sulphate,  $Ce_2(SO_1)_2$  (NII<sub>1</sub>)<sub>2</sub>SO<sub>1</sub> 8H,O, is a monochine salt (see p. 263) of density 2.523. It loses 6H<sub>2</sub>O at 100′, and the remaining 2H<sub>2</sub>O at 150°. The solubility, in grams of  $Ce_2(SO_1)_4$  (NII<sub>1</sub>)<sub>2</sub>SO<sub>1</sub> per 100 grams of water, is as follows (Wolff) - 2

Above 45' the anhydrous double salt is the stable phase in contact with the solution, and its solubility is as follows (Wolff) .-

The sulphates of hydrazine and hydroxylamine combine with cerous sulphate 5

Cerous thallous sulphate, Ce (SO<sub>1</sub>)<sub>3</sub> Tl<sub>2</sub>SO<sub>1</sub> 4H<sub>2</sub>O, erystallises in monoclime prisms (a=b=c=1 1309=1=0 7059,  $\beta$ =91° 53′). The salt Ce<sub>2</sub>(SO<sub>1</sub>)<sub>4</sub> 3Tl<sub>2</sub>SO<sub>4</sub>.  $\hat{\mathbf{H}}_{2}\mathbf{O}$  has also been described

Cerous cadmium sulphate, Ce<sub>2</sub>(SO<sub>1</sub>), CdSO<sub>1</sub> 6H<sub>2</sub>O, crystallises in the orthorhombic system (a b c = 1 1336 , 1 0 7535).

Cerous nutratosulphate, Ce(NO<sub>1</sub>)(SO<sub>1</sub>) H<sub>1</sub>O, crystallises out when a solution of cerous sulphate in concentrated intricated is evaporated on the steam-bath.

Cerous dithionate,  $(e_2(8_20_b), 2411_20)$ , is extremely soluble in water (Jolin). The hydrate with  $15H_20$  separates from solution at  $15-17^\circ$  in triclinic crystals (a  $b \cdot c = 0.5917 - 1.1.1912$ ,  $a = 81^\circ 26'$ ,  $\beta = 105^\circ 21'$ ,  $\gamma = 86^\circ 38'$ ) of density 2 288, the hydrate with 3H2O separates above 20', in triclinic

Barre, loc, et ... Solid phases not specified
 Wolff, Zetts h. anorg. Chem., 1905, 45, 89
 Kolb, Zettsch. anorg. Chem., 1908, 60, 123
 Wyrouboff, Bull. Soc. franc. Min. 1891, 14, 83
 Wyrouboff, loc, ett
 Wyrouboff and Verneult, Ann. Chem. Phys., 1906, [viii.], 9, 290

orystals (a: b: c=0.5807:1:1.2030; a=89° 25′,  $\beta$  = 96° 9′,  $\gamma$  = 96° 18′) of density 2.631. A hydrate with 12H<sub>2</sub>O is also known <sup>2</sup>

Cerous selenite, Ce<sub>2</sub>(SeO<sub>3</sub>), 12H<sub>2</sub>O, is precipitated from cerous acctate and selenious acid, and loses 9H<sub>2</sub>O over sulphuric acid. The acid selenites, Ce<sub>2</sub>(SeO<sub>3</sub>)<sub>3</sub>, H<sub>2</sub>SeO<sub>3</sub>, 4H<sub>2</sub>O (or 5H<sub>2</sub>O) and Co<sub>2</sub>(SeO<sub>3</sub>)<sub>3</sub>, 3H<sub>2</sub>SeO<sub>3</sub>, 2H<sub>2</sub>O, have also been described.8

Cerous selenate, Ce<sub>2</sub>(SeO<sub>1</sub>), and hydrates with 4, 5, 7, 8, 10, 11, and 12H<sub>2</sub>O have been described by Cingolani, while John has described hydrates with 6 and 9H2O. The hydrates are less soluble in hot than in cold water.4 The following double salts have been prepared by John -

$$\begin{array}{l} \operatorname{Cc_2(SeO_1)_3} \operatorname{5K_2SeO_1} \\ \operatorname{Ce_2(SeO_1)_3.Na_2SeO_1} \operatorname{5H_2O} \end{array}$$

. : 4

Cerous chromate -Potassium chromate or dichromate gives with a solution of a cerous salt a yellow precipitate of a basic salt?

Cerous molybdate, Ce<sub>2</sub>(MoO<sub>1</sub>), forms yellow crystals of density 4 56.6 Cerous tungstate, Ce<sub>2</sub>(WO<sub>1</sub>), forms sulphur-yellow crystals of density 6:514 and specific heat 0 0821.7 It melts at 1089° 8 For paratungstates, double and complex tungstates, see p. 265.

Cerous silicotungstate, Co<sub>4</sub>(W<sub>12</sub>SiO<sub>10</sub>)<sub>3</sub>. See p. 266.

### CERIUM AND THE NITROGEN GROUP

Cerous nitride, CeN.—The preparation of this compound has been described (p 267). Gas is violently evolved when a little water is added to the intride .

$$CeN + 2H_2O = CeO_2 + NH_3 + H.$$

It dissolves in acids, producing cerous and ammonium salts, eg.:-

$$2\text{CeN} + 4\text{H}_2\text{SO}_4 = \text{Ce}_2(\text{SO}_4)_3 + (\text{NH}_1)_2\text{SO}_4$$

Cerous azide (hydrazoate, trinitrule) is an explosive salt 9

Cerous nitrite, Ce(NO<sub>2</sub>)<sub>3</sub> xll<sub>2</sub>O, is extremely soluble in water, from which it separates in transparent, amber yellow crystals. The solid salt is very unstable, and its aqueous solution also readily decomposes when warmed,

so that the salt has not been obtained in a state of purity. O

Cerous nitrate, Cc(NO<sub>1</sub>), 6H<sub>2</sub>O, forms very deliquescent, triclinic

crystals. A solution of the salt may be prepared from cerous sulphate-and barium nitrate, by heating ceric hydroxide or oxide with nitric acid and a reducing agent, eg hydrogen peroxide, or by heating corous evalute with

Wytouboff, Bull Soc fram; Min , 1891, 14, 83
 Morgan and Cahen, Trans Chem. Soc., 1907, 91, 475
 John, Ioc. ett; Nilson, Nova Acta Soc. Upsala, 1875, [111], 9, No. 7, Her., 1875, 8, 655; Bull. Soc. chim , 1875, [1], 23, 494.
 Cingolani, Gazzetta, 1908, 38, 1 292; Atti R. Accad Lincei, 1908, [v.], 17, i. 254; John Low ett.

<sup>4</sup> Cingolain, Gazzetta, 1908, 38, 1 292; Atti R. Accad Lineal Jolin, loc. cit.
b Boringer, Annalen, 1842, 42, 138.
6 See p. 265; also Didier, Compt. rend., 1886, 102, 823
7 See p. 265, also Didier, loc. cit.
b Zambonini, Atti R. Accad Lineal, 1913, [v.], 22, 1. 519
Cuttus and Darapsky, J. prakt. Chem., 1900, [ii], 61, 408
10 Morgan and Cahen, Trans. Chim. Soc., 1907, 91, 475.
11 Mangnao, Ann. Chim. Phys., 1873, [v.], 30, 56. See p. 268.

concentrated nitric acid.1 The crystals lose 3H,O at 100°, and begin to decompose at 200°.

Cerous ammonium nutrate, Ce(NO<sub>3</sub>), 2(NII<sub>4</sub>)NO<sub>3</sub> 4H<sub>2</sub>O<sub>5</sub>, described on p. 268, has been prepared by numerous chemists.<sup>2</sup> Its solubility, in grams of Ce(NO<sub>3</sub>)<sub>3</sub> 2(NII<sub>1</sub>)NO<sub>3</sub> per 100 of water, is as follows (Wolff):-

65·1°

The salt 2Ce(NO<sub>3</sub>)<sub>3</sub> 3(NH<sub>4</sub>)NO<sub>3</sub> 12H<sub>5</sub>O separates from aqueous solution of 0°.4

Cerous cossum nutrate,  $Ce(NO_3)_3$  2CsNO<sub>3</sub> 2H<sub>2</sub>O, crystallises in the monochine system (a b c=1 2052 1 0 9816 ,  $\beta$  103 41).

Cerous rubidium nitrate, Cc(NO<sub>2</sub>), 2RbNO<sub>3</sub>4H<sub>2</sub>O, has been described

Cerous potassium nitrate, Ce(NO<sub>3</sub>)<sub>3</sub> 2NAO 2H<sub>3</sub>O, crystallises in the orthorhombic system (hemimorphic, a b e 0.5227 1 0.5704) 5

Cerous sodium nitrate, Ce(NO<sub>3</sub>)<sub>3</sub> 2NaAO<sub>3</sub>H<sub>2</sub>O, forms hygroscopic crystals

of density 2 65 at 0° 6

Circus thallous nitrate, Ce(NO3), 2TINO , 4H2O, has been described (p. 268). The double salts of cerous mitrate and the mitrates of bivalent metals, 2Ce(NO<sub>3</sub>)<sub>3</sub>3M<sup>II</sup>(NO<sub>3</sub>)<sub>2</sub>,24H<sub>2</sub>O, have been already described <sup>7</sup>

Cerous hypophosphite, Ce(H2PO2), H2O, crystallises in thin, prismatio crystals s

Cerous orthophosphate, CePO1 -- The dihydrate, CePO1 2H2O, is obtained by double decomposition between a cerous salt and phosphoric acid or an alkali phosphate " When dried it forms a white, amorphous powder, from which orthorhombic crystals of density 5 09, closely resembling the mineral monante, may be obtained by heating to redness with excess of fused cerous chloride. The phosphate dissolves in the molten chloride and crystallises out on cooling.10

The double phosphates, K,PO, 2CePO, and Na,PO, 2CePO, have been obtained by Ouvrard.

Cerous pyrophosphate,  $Ce_1(P_2O_7)$ ,  $12H_2O$ , cerous hydrogen pyrophosphate,  $CoIIP_2O_7$ , and cerous sodium pyrophosphate,  $CeNaP_2O_7$ , have been described, and likewise eerous metaphosphate,  $Ce(PO_3)_3^{-11}$ . Cerous pyrophosphate is readily soluble in dilute mineral acids.

John, Bull Soc chim, 1871, [n.], 21, 533, Lange, J prakt Chem, 1861. 82, 129.
 Mangnac, low ed., Fock, Zetsch Kryst Min, 1894, 22, 37, Kraus, chid, 1901, 34,
 Wolff, Zetsch anory. Chem, 1905, 45, 89.
 Holzmann, J prakt. Chem, 1861, 84, 76.
 Wyronboff, Bull. Soc franç. Min., 1907, 30, 299, Jantsch and Wigdorow. Zeitsch. anory. Chem., 1911, 69, 221.
 Lange, loc. ett., Fock, loc. ett., Wyronboff, loc. ett.
 Jantsch and Wigdorow, loc. ett.
 See p. 299, 298 bl. lleymann. loc. ett.
 See p. 299

Jantsch and Wigdorow, loc. ett.
 See p. 269, also Holzmann, loc. ett.; Lange, loc. ett., Z-chresche, J. prakt. Chem.,
 1869, 107, 65, Rammelsberg, Pogg. Annalen, 1859, 108, 435
 Rammelsberg, Ber., 1872, 5, 492.
 John, loc. ett., Hartley, Trans. Chem. Soc., 1882, 41, 202
 Radominski, Compt. rend., 1875, 80, 301
 See also Ouverd., third., 1868, 107, 37;
 Grandeau, Ann. Chim. Phys., 1886, [vi.], 8, 193, Holm, Interqueal Insortation (Munich, 1902)

<sup>11</sup> Rosenheim and Triantaphyllides, Ber., 1915, 48, 552 (Cc<sub>1</sub>(P<sub>2</sub>O<sub>7</sub>), 12H<sub>2</sub>O), John, loc. it (CeHP<sub>2</sub>O<sub>7</sub>), Wallroth, Bull. Soc. chim., 1883, [n.], \$9, 316 (CcNaP<sub>2</sub>O<sub>7</sub>), Rammelsberg, 12 loc. cit. (Ce(PO<sub>2</sub>)<sub>2</sub>). See also Johnston, Ber., 1889, 22, \$76.

Cerous vanadate, CeVO, may be obtained in dark-red, dichroic needles by fusing sodium orthovanadate with corous chloride. The compound  $\text{Ce}_2\text{O}_3.5\text{V}_2\text{O}_5$  27H<sub>2</sub>O is also known.

# CERIUM AND THE CARBON GROUP.

Cerium carbide, CeC, is prepared by reducing ceric oxide with carbon in the electric furnace. It has been already described. The reduction of ceria by carbon probably proceeds in three stages :-

$$\begin{array}{ll} 2 {\rm CeO}_2 + {\rm C} &=& {\rm Ce}_2 {\rm O}_3 + {\rm CO} \\ {\rm Ce}_2 {\rm O}_3 + 9 {\rm C} &=& 2 {\rm CeC}_3 + 3 {\rm CO} \\ {\rm CeC}_3 &=& {\rm CeC}_2 + {\rm C} \end{array}$$

and red crystals of the carbide CeC, may be isolated under suitable conditions. They are decomposed by water with the evolution of acetylene —

$$2\text{CeC}_3 + 6\text{H}_2\text{O} - 2\text{Ce(OII)}_3 + 3\text{C}_2\text{H}_2$$
.

Cerium silicide, CeSi, is propared by heating cene oxide with silicon in the electric furnace. It forms small, lustrous crystals of density 5.67. It is decomposed by fluorine in the cold and by the other halogens when heated. It burns in oxygen at a red heat. Unlike the carbide, ceruum silicide is scarcely affected by water or dilute alkalies, even when boiling; and although it dissolves in dilute hydrochloric and sulphuric acids, the gas evolved is pure hydrogen 3

Cerous carbonate,  $\text{Ce}_2(\text{CO}_3)_3$  5H  $_3\text{O}_7$  is known, 4 and also the following double carbonates -5

$$\begin{array}{lll} \text{Ce}_2(\text{CO}_3)_3 \text{ K}_2(\text{CO}_3 \text{3H}_2 \text{O} \text{ (and } 12\text{H}_2 \text{O}) & 2\text{Ce}_2(\text{CO}_3)_3 \text{3Na}_2 \text{CO}_1 24 \overset{?}{\text{H}}_2 \text{O} \\ \text{Ce}_2(\text{CO}_3)_3 \text{2Na}_2 \text{CO}_3 2\text{H}_2 \text{O} & \text{Ce}_3(\text{CO}_3)_3 (\text{NH}_4)_2 \text{CO}_1 6\text{H}_2 \text{O} \end{array}$$

For preparation and properties, see p. 271.

Cerous thiocyanate, Ce(CNS), 71120, and the double salt, Ce(CNS),

SHg(CN)<sub>2</sub>,12H<sub>2</sub>O, are known. (John, see p. 272.)

Cerous platinocyanide, 2Ce(CN)<sub>3</sub>,3Pt(CN)<sub>2</sub>,18H<sub>2</sub>O, crystallises in yellow prisms (sp. gr. 2.657) having a blue reflex 6

Cerous ethylsulphate, Ce(C<sub>2</sub>H, SO<sub>1</sub>)<sub>1</sub>, 9H<sub>2</sub>O, has been described (p. 278).

Cerous acetylacetonate, Ce(CH<sub>3</sub> CO.CH CO.CH<sub>1</sub>)<sub>3</sub>, has also been described (p. 278). scribed (p. 279)

Cerous oxalate, Ce<sub>2</sub>(C<sub>2</sub>O<sub>1</sub>), 10H<sub>2</sub>O, may readily be obtained by double decomposition as a white, crystalline powder. The oxalate when air-dried has the above composition, but crystalline hydrates containing 311,0, 9H,0,

See p 270; also Sterba, Compt. rend., 1902, 134, 1056, Ann Chim. Phys., 1904, [vii.], 2, 223
 Dannens, Compt. rend., 1913, 157, 335.
 Sterba, Compt. rend., 1902, 135, 170, Ann. Chim. Phys., 1904, [viii.], 2, 229.
 Beninger, Annalen, 1842, 42, 138, John, Bull. Soc. chim., 1871, [ii.], 21, 533; Czudnowicz, J. prakt. Chem., 1861, 82, 277
 John, loc. cit.; R. J. Meyer, Zeitsch. anorg Chem., 1904, 41, 97; K. A. Hofmann and Hoschele, Ber., 1915, 48, 20 (Co - NH<sub>4</sub>).
 Czudnowicz, J. prakt. Cheme, 1860, 80, 16, Lange, ibid., 1861, 82, 129. See also p. 272.

also p. 272.

and 11H<sub>2</sub>O have also been prepared. For the solubility of cerous exalate in water and acids, see p. 273

Cerous oxalate is transformed into a crystalline oxalochloride when it is dissolved in hot, concentrated hydrochloric acid, and the solution allowed to cool 2

Cerous salts of other organic acids - The following salts have been described -

Cerous formate, \$7 acetate, \$3.4.5.6 propronate, butyrate, 6 isobutyrate, 6 monochlor, 6 dichlor-17 and trichlor-acetate, 6 succinate, 5 citiate, 5 11 tartrate, 5 11, 11 ylycollate, 12, 15 dimethylphosphate, 10 crotonate, 12 malate 12 and acid malate, 12 maleate 12 and acid malate, 12 maleate 12 and acid malate, 12 fumarate 12 and acid fumarate, 12 malonate 11 12, 11 and acid and acid materite, 12 fumarate 12 and acid fumarate, 12 malonate 11 12, 11 and acid malonate, 12 benzoate, 5 salicylate, 7 phthalate 12 and acid phthalate, 12 hippurate, 5 phenos yacetate, 8 sulphanilate, 17 ben encsulphonate, 17 m-nitrohen encsulphonate, 14 p-dibromobenzenesulphonate, 16 1 1 2-bromonitrohen encsulphonate, 9 phenol-psulphonate, 17 naphthionate, 17 happithol-6-sulphonate, 17 camphon \( \beta\) sulphonate, 18 anthraquinonesulphonate 18 and numerous naphthalenesulphonate 18

Cerous silicate, Ce2(SiO3), may be obtained in prismatic crystals by fusing together cerous chloride or oxychloride, silica and sodiu n or calcium chloride 19

CERIUM AND BORON.

Cerous metaborate, Cc(BO,),, has been described 20

# CERIC COMPOUNDS.21

Ceric salts are derived from the feebly basic oxide CeO2, and are yellow, orange, or red in colour. Being salts derived from a weak base, they are considerably hydrolysed in aqueous solution, and normal cene salts of weak acids are not known. Further, the normal chloride and intrate are only known in combination as double, or possibly complex, salts Aqueons solutions

of ceric salts, owing to hydrolysis, react strongly acid. The solutions are very unstable and are easily reduced to the cerous state. On the ionic hypothesis, the instability is attributed to the ceric ion, Ce''', the oxidation potential of which is greater than that of ovygen; acid solutions of ceric salts accordingly behave as if they were supersaturated with oxygen. In alkaline media, however, ceric compounds are not readily reduced to the cerous state, while the converse change is very easy to effect

Hydrolysis of Ceric Salts.—The hydrolysis of a ceric salt in cold, aqueous solution proceeds for a long time after the solution has been prepared, and the colour fades away very perceptibly, the hydrolysis may be hastened and increased by raising the temperature. A freshly prepared solution of a ceric salt immediately darkens in colour when a inneral acid is added, since the degree of hydrolysis is thereby diminished, with an old solution, however, the deepening of the colour takes place very slowly. Again, a fresh solution is immediately decolorised by hydrogen peroude, with the formation of a cerous salt and oxygen, but an old solution first turns dark red in colour when similarly treated, and is only slowly reduced <sup>2</sup> By the hydrolysis of ceric salts basic ceric salts are produced, which, under suitable circumstances, may be utilised for the separation of ceria from the other rate earths (see p. 332).

Colloidal Ceric Compounds.—A cold solution of ceric nitrate in which hydrolysis has proceeded to a considerable extent, either by long standing or by heating, contains a colloidal hydrosol, which coagulates when intric acid is added and is almost quantitatively precipitated if 12 cubic centimetres of concentrated intric acid are added for every 100 cubic centimetres of dilute ceric solution present. The hydrogel thus obtained, when dired over potassium hydroxide, forms an amber-coloured, horny, translucent solid of the composition 4CcO<sub>2</sub> N<sub>2</sub>O<sub>3</sub> 5H<sub>2</sub>O. It easily changes back into the hydrosol in contact with water, giving a greenish, limpid solution unless it is very concentrated, when a faint opalescence is observable. Submitted to dualysis, the solution loses intric acid; the whole of the acid present in the hydrosol, however, cannot be thus removed, the decomposition ceasing when the ratio 28CeO<sub>2</sub> 1N<sub>2</sub>O<sub>5</sub> is reached

When the solution of the basic nitrate hydrosol is treated with one-fifth its volume of concentrated hydrochloric acid, the cerum is almost quantitatively precipitated as a hydrogel which has the composition  $4\text{CeO}_2$ .2HCl.  $34\text{H}_2\text{O}$  and is very similar in properties to the basic intrate; in solutions of both of these compounds one-half of the acid present may be neutralised with sodium hydroxide before ceric hydroxide begins to be precipitated. When, however, a dilute solution of a dibasic acid (or, better, its ammonium salt) is added to the basic nitrate hydrosol, a hydrogel is precipitated which does not dissolve in water. The basic sulphate,  $4\text{CeO}_2.6\text{O}_3.5\text{H}_2\text{O}$ , for instance, which resembles the chloride and nitrate in appearance, loses half its sulphuric acid when washed with warm water, but does not dissolve to any appreciable extent.

The addition of ammonia to the basic nitrate hydrogel converts the latter into a hydroxide which has the composition  $8\text{CeO}_2.11\text{H}_2\text{O}$  when dried over potassium hydroxide, and the same horny appearance as the basic nitrate.

Colloidal compounds similar in properties to the preceding may be prepared containing lanthanum, prascodymium, etc., in addition to cerium.

<sup>1</sup> Baur and Glaessner, Zeitsch. Et trochem., 1903, 9, 534.
R. J. Meyer and Jacoby, Ber., 1900, 33, 2135; Zeitsch. anorg. Chem., 1901, 27, 359.

Concerning the constitution of these colloidal substances, little can be said beyond the statement that they do not appear to be basic salts of the ordinary type. Wyrouboff and Verneuil regard them as derivatives of polymerised or "condensed" ceric hydroxides, such as  $\text{Ce}_{21}\text{O}_{15}(\text{OH})_{60}(\text{OH})_{60}$  in which only part of the hydroxyl (the  $(\text{OH})_{60}$  of the preceding formula, for example) is capable of reacting with acids, they speak of the compounds as being derived from certain metaoxides or mixed metaoxides, the term "meta-" being used as in naming "condensed" acids, e.g. metastannic and metatungstic acids. There is very little evidence, however, for the molecular formulæ ascribed to these

compounds by the French chemists.2 Colloidal basic ceric compounds of another type have also been prepared by Wyrouboff and Verneuil, who speak of them as derivatives of a paraoxide. A description of the basic nitrate may be given. When cerous oxalate is calcined in air at the lowest possible temperature, a canary yellow residue of ceria is obtained containing 2.9 per cent of water. It is quite indifferent towards concentrated intric acid, but is transformed into a white, gelatinous substance when heated to 100° with 3 per cent intric acid for several hours. This substance, when separated from the dilute acid by decantation, may be dissolved in water. The solution has a decidedly milky appearance, the hydrosol it contains may be completely precipitated by the addition of intric acid (2 per cent ) or ainmonium nitrate, and dries at 100° to a very palecoloured, horny, translucent mass, soluble in water Wyrouboff and Verneuil propose the molecular formula  $20{\rm CeO_2~N_2O}$ ,  $5{\rm H_2O}$  for the substance, basic chlorides, sulphates, etc., and another modification of ceric hydroxide may be prepared from it as in the case of the meta nitrate Moreover, the French chemists state that when cerons hydroxide, precipitated from a cerous salt by means of ammonia, is oxidised to cerie hydroxide by a current of air, the product is almost entirely insoluble in boiling concentrated intric acid, and that the insoluble portion is colloidal, dissolving in water to form a solution of the para-hydroxide

Further work upon these interesting substances is very desirable

Conversion of Cerous into Ceric Compounds - Owing to the great instability of ceric chloride, this transformation cannot be effected in solutions acidified with hydrochloric acid

(1) Oxidation in nitric acid solution - A solution of cerous nitrate in concentrated intric acid may be oxidised to ceric intrate to the extent of 6-8 per cent. by evaporation at 100° 3. In the presence of the requisite amount of alkalı nitrate, some 30 per cent or more of the cerous salt may be exidised, a result that is attributed to the transformation of the ceric ions Ce''', as they are produced, into the complex ion Ce(NO3),", the probable existence of which has been shown by Meyer and Jacoby (see p 400)4 Provided that (i.) a cerous salt is readily soluble, and the corresponding ceric salt only sparingly soluble in concentrated intric acid, and (ii ) the acid from which the cerous salt is derived is not attacked by intrie acid, eg -iodu, phosphoric, and arsenic acids, the oxidation of the cerons to the ceric salt may be readily accomplished by boiling it with concentrated nitric acid.5

It would be more correct to say metahydroxides.

<sup>&</sup>lt;sup>2</sup> Wyrouhoff and Verneul, Compt. 1898, 127, 863; 1849, 128, 501. Bull. Soc. chm, 1899, [ni.], 21, 118; Ann Chun Phys., 1906, [vii.], 9, 289.

<sup>3</sup> Giles, Chem. News, 1905, 92, 1; Bathani, Att. R. Acc. al. Lincei, 1907, [v.], 16, i. 395.

<sup>4</sup> Parly of the Compt. 1895, 92, 1; Bathani, Att. R. Acc. al. Lincei, 1907, [v.], 16, i. 395.

<sup>5</sup> Barbieri, 181d., 1907, [v.], 16, 1. 644, Ber, 1910, 43, 2214.

Oxidation in nitric acid solution may be effected by the use of various oxidising agents, ey lead peroxide and bismuth tetroxide, the processes are useful in connection with the analytical chemistry of cerium 1

(ii.) Oxidation in sulphuric acid solution - This may be accomplished by means of ammonium persulphate or sodium bismuthate. The processes are useful for analytical purposes (see p 373), and the former may also be utilised for the preparation of pure ceria (see p 333).

(iii.) Electrolytic oxidation - The electrolysis of cerous intrate or sulphate between platinum electrodes in neutral or slightly acid solution leads to the separation of ceric hydroxide or basic ceric salt at the anode, but in the presence of sufficient mineral acid, cene salt is produced in solution. As much as 95 per cent of the cerum may be oxidised under suitable conditions 2

(iv.) Oxidation in alkaline media.—Cerous hydroxide is readily oxidised to ceric hydroxide by alkali hypochlorite or hypobromite. Cerous salts are also readily converted into ceric hydroxide by potassium permanganate in the presence of a base, e.g. sodium hydroxide or magnesia. These reactions are of considerable value both in preparation work (see pp. 335, 336) and in analytical chemistry (see p 373) A solution of corous carbonate in potassium carbonate is easily oxidised by oxygen or hydrogen peroxide (see p. 401)

In alkaline or acetic acid solution, cerous compounds may be oxidised to

perceric compounds by means of hydrogen peroxide (see pp. 401-3).

Conversion of Ceric into Cerous Compounds—In acid solution ceric salts may be reduced to the cerous state with great ease by numerous reducing agents, e.g. hydrogen peroxide, sulphurous acid, hydrochloric, hydrobiomic, and hydriodic acids, oxalic acid, stannous chloride, ferrous sulphate, etc The neatest method is that involving the use of hydrogen peroxide, but on a large scale it is rather expensive. The transformation from ceric nitrate or sulphate into cerous ovalate is readily effected in warm, acid solution by the addition of oxalic acid

The conversion of corrum dioxide into cerous salts is worthy of special notice, inasmuch as the dioxide is insoluble in hot, concentrated hydrochloric or nitric acid The conversion into cerous intrate may be accomplished very neatly by warming the oxide with moderately concentrated intric acid and adding hydrogen perovide from time to time. The conversion into cerons sulphate or chloride may be brought about by heating the oxide with hydroquinone and an excess of the requisite acid in aqueous solution, the hydroquinone being converted into benzoquinone and quinhydrone 3. Ceria may also be converted into cerous sulphate by heating it with concentrated sulphuric acid until it has been converted into cenic sulphate and reducing its aqueous solution with sulphurous acid, while it may be converted into anhydrous cerous chloride by heating in the vapour of disulphur dichloride (p. 252), and into a solution of cerous chloride (plus alkali chloride) by heating with concentrated hydrochloric acid and an alkali iodide (p. 332).

See p. 373. For the use of the lead peroxide method in preparing pure ceria, see Robinson, Proc. Roy. Soc., 1884, 37, 150, Chem. News, 1884, 50, 251
 Eik, Jahreber, 1870, p. 319, Smith, Ber., 1880, 13, 754, Bircout, Compt. rend., 1894, 118, 145; Kolle, Inauqueral Dissertation (Zurich, 1898), von Knone, Zeitsch. angen. Chem., 1897, 10, 685, 717, Joh, Compt. rend., 1899, 128, 101; Sterba, ibid., 1901, 133, 221, Ann. 5(Chim. Phys., 1904, [viii.], 2, 193; Muhlbach, Inauqueral Dissertation (Munich, Tech. Hochschule, 1903), Killitschewski, Inauqueral Dissertation (Gessen, 1904), Plancher and Barbieri, Atti R. Accad. Lince, 1904,[v.], 14, i. 119. See also p. 334.
 Marino, Gazzetta, 1907, 37, 1. 57.

## CERIUM AND THE FLUORINE GROUP.

Ceric fluoride, CeF4.II<sub>2</sub>O, is obtained as a brown powder, insoluble in water, by acting upon cene hydroxide with hydrogen fluoride and drying the residue at 100°. When heated, water, hydrogen fluoride, and possibly fluorine, are evolved 1 According to Brauner, the double salt 2CeF, 3KF 2H2O is produced by the action of potassium hydrogen fluoride on ceric hydroxide. Rimbach and Kihan 2 could not obtain this salt, but have prepared a series of double salts of the type  $2\text{CeF}_4$  MF<sub>2</sub>.7H<sub>2</sub>O, where M = Cd, Cu, Co, Ni, or Mn, They are decomposed by water

Ceric chloride, CeCl, -This salt has not been isolated. It can be obtained in solution, but the solution cannot be kept without decomposing. Thus, cene hydroxide dissolves in cold concentrated hydrochloric acid with the production of a dark red solution, chlorine, however, is slowly evolved, and ultimately a solution of cerous chloride remains. The decomposition

proceeds rapidly in hot solutions

By preparing a solution of ceric chloride in methylalcoholic hydrogen chloride and adding an organic base, crystalline double chlorides may be prepared. The pyridine, quinoline, and triethylamine salts have thus been made, and ascribed the formula CeCl  $_1$  2(C, H, N HCl), CeCl  $_1$  2(C, H, N HCl), and CeCl<sub>1</sub> 2[(C<sub>2</sub>H<sub>5</sub>)<sub>3</sub>N HCl] The compositions of these compounds, however, vary with the amounts of the reacting salts 1

Ceric iodate, Cc(10,)1, may be prepared by heating together cerous intrate, iodic acid, and concentrated intric acid. It is a yellow crystalline salt, very sparingly soluble even in boiling intric acid, and hydrolysed somewhat by water 5

## CERIUM AND THE OXYGEN GROUP,

Ceroceric oxide,  $Ce_1O_7$  or  $Ce_7O_{19}(l)$  —When certain dioxide is heated to a red heat in a current of dry hydrogen free from air, it is partially reduced and loses in weight by about 2 per cent. The reduction proceeds most rapidly with an oxide that has not previously been intensely ignited, but such an oxide is hable to have retained a little water, accordingly it is difficult to obtain reliable analytical data concerning the product formed by the reduction. The lower oxide is considered to correspond to the violet céroceric hydroxide (described below) and to the acid salt known as cerocerie hydrosulphate (p. 397), hence Meyer regards it as Ce<sub>1</sub>O<sub>2</sub>, and Wyrouboff and Verneuil as Ce<sub>7</sub>O<sub>12</sub> It is a blue or bluish-black solid which readily absorbs oxygen and is sometimes pyrophoric 6

Ceroceric hydroxide, Ce(OH), Ce(OH), or 1Ce(OH), 3Ce(OH), (3).— When white cerous hydroxide is exposed to an or oxygen, it absorbs oxygen and is eventually converted into yellow cene hydroxide, but the initial product

<sup>1</sup> Brauner, Trans Chem Soc., 1882, 41, 68
2 Rumbach and Kihan, Annalen, 1909, 368, 101.
3 Koppel, Zeitsch. anorg Chem. 1898, 18, 305
4 Grant and James, J. Amer. Chem. Soc., 1915, 37, 2652.
5 Batberi, Allt. R. Accad Linect, 1907, [v.], 16, 1-614.
6 Stetba, Compt. rend., 1901, 133, 221, Ann Chem. Phys., 1304, [viii.], 2, 210; Br. J. Meyer, Zeitsch, anorg Chem., 1903, 37, 378. Wytoutoff and Verneul, Ann Chim. Phys., 1906, [viii.], 9, 289 The sarly workers obtained erroneous results, see Beringer, Annalen, 1842, 42, 138, Bunsen, ibid., 1858, 105, 40, 45; Rammelsberg, Popy Annalen, 1859, 108, 40, Winkler, Bec., 1891, 24, 873.

of oxidation has a violet or purple tint. A precipitate of the same violet colour is obtained by adding an alkali hydroxide to a solution of a mixture of a cerous and a ceric salt, and the colour attains a maximum intensity when the ratio Ce<sup>III</sup> Ce<sup>IV</sup> has a certain definite value, which, however, can only be approximately estimated. Wyrouboff and Verneuil consider that this intermediate hydroxide is obtained in a pure state by precipitating coroceric hydrosulphate with alkalı, and, if this is the case, its composition is probably expressed by one or other of the alternative formulæ given above. The violet hydroxide becomes bluish-black in colour when dried in vacuo, but a slight oxidation to ceria is always observed.1

Ceric oxide, cerium dioxide, or ceria, CeO2, may be prepared by the ignition of cerous or ceric hydroxide, nitrate, sulphate, etc., or by the ignition of the cerous salt of any volatile oxyacid; it is perhaps most commonly prepared by the ignition of cerous oxalate

Cena is an amorphous powder of specific gravity 6 405 at 17° when prepared from the oxalate, and 6 99 when obtained from the nitrate (Sterba); its specific heat is 0.0877 (0°-100°).2 There has been a great deal of discussion over the question of the colour of ceria. It might be anticipated that ceria, like zirconia and thoria, would be white. Most experimenters agree that pure ceria 3 has a pale yellow colour The depth of colour depends upon the temperature at which the oxide has been calcined and the salt from which it has been prepared. When prepared by the prolonged ignition of cerous sulphate at a white heat, its tint is so slight that it may almost be said that the ceria is white,4 but when obtained at a lower temperature by the ignition of cerous oxalate or ceric ammonium intrate it has a more pronounced tint, usually described as that of pale chamois. On the other hand, Spencer claims that when ceric sulphate is heated for a prolonged period at temperatures below a red heat, cerium dioxide is formed, which is pure white, further, that when the cena is heated above a red heat it shrinks in volume and becomes pale yellow in colour Ceria darkens in colour very markedly when heated, but returns (practically) to its original colour when cooled 5 ceria is contaminated with a little of the other earths of the cerium group, it is salmon-coloured, reddish-brown, or brown, according to the extent of contamination and the temperature of ignition. The coloration is attributed mainly to the presence of prascodymia, or rather, its peroxide.

<sup>&</sup>lt;sup>1</sup> Dennis and Magee, J. Amer. Chem. Soc., 1894, 16, 662, Wyrouboff and Verneuil, loc. ctt.; Rammelsheig, Pogg. Annalen, 1859, 108, 45

<sup>2</sup> Nilson and Pettersson, Compt. rend., 1880, 91, 232; Ber., 1880, 13, 1159.

<sup>3</sup> For the preparation of which see p. 337

<sup>4</sup> In this sense of the term, certa is said to be white by Wyrouboff and Verneuil (Compt. 1002) and 1002 for the control of the term, certa is said to be white by Wyrouboff and Verneuil (Compt. 1002) and 1002 for the control of the term, certa is said to be white by Wyrouboff and Verneuil (Compt. 1002) and 1002 for the control of the term, certa is said to be white by Wyrouboff and Verneuile that is the control of the term, certa is said to be white by Wyrouboff and Verneuile that is the control of the term, certain the control of the term of

an this sense of the term, certa is said to be white by wyrothom and verneth (Compt. rend., 1897, 124, 1800; Ann Chim. Phys., 1906, [vin ], 9, 356), who consider that it is possible to distinguish different kinds of white, and that the colour of cens prepared from cerous sulphate by ignition is not definite enough to be described as anything but a particular kind of white.

cular kind of white.

\* Steba (Compt. rend., 1901, 133, 221; Ann. Chim. Phys., 1904, [viu.], 2, 193) has prepared ceria devoid of any yellow tint, but Brauner has shown that the ceria so prepared contains traces of impurities. Other chemists (e.g. Wolf. Amer J. Sci., 1868, [ii.], 46, 54; Moissan, Compt. rend., 1897, 124, 1238) have also described ceria as being white. Steba, after preparing ceria in various ways, concluded that it could be white or pale cition yellow in colour, and that the yellow colour, which could not be attributed to non, platinum, or other rare earth clements, might possibly be due to the presence of traces of a higher oxide than the dioxide Spencer's results appear to lend support to this riew. On the question of the colour of pine ceria, see Sterba, loc. cit; Wylouboff and Verneuil, loc. cit; R. J. Møyer, Zelisch anorg. Chem., 1903, 37, 378; Brauner, chul., 1903, 34, 207; Neish, J. Amer. Chem. Soc., 1909, 31, 517; J. F. Spencer, Trans. Chem. Soc., 1915, 107, 1285.

Ceria is readily obtained in the crystalline form by adding anhydrous cerous sulphate to molten magnesium chloride, allowing to cool slowly, and extracting the mass with hydrochloric acid. The crystals belong to the regular system and exhibit faces of the cube and octahedron. They are practically colourless, very hard and very brilliant, the refractive index being high (about 1.9), they have a density of 7.3.1 If small quantities of another rare earth sulphate are added in the preparation, e.g. neodymium, praseodymium or erbium sulphate, beautiful coloured crystals may be obtained 2

Ceria does not melt at c 1900°, but it volatilises in vacuo quite rapidly at that temperature 3 It readily melts in the electric furnace and attacks the containing vessel

Ceria is a very stable oxide, but it can be reduced to the metallic state by heating it with aluminium 1 or magnesium. 5 By neither of these methods, however, has a regulus of the metal been obtained The action of other reducing agents is discussed in connection with cerous oxide and ceroceric oxide

Crystalline cena is very resistant towards acids and alkalis. The amorphous substance after ignition is insoluble in hydrochloric or intric acid, except in the presence of a suitable reducing agent, e.g. hydrogen peroxide, hydrodic acid, or stannous chloride, when it passes into solution as a cerous salt. Concentrated sulphuric acid converts it into ceric sulphate, while moderately concentrated and causes partial reduction to cerous sulphate and dilute and has no perceptible action

Ceria acts as an oxygen-carrier towards other substances, in a manner that is not at present understood 6 It may therefore be employed as the catalyst in Dennstedt's method for the combustion of organic compounds 7

Ceria is a very weakly basic oxide, it is possible that it can also act as a feebly acidic oxide. It has been pointed out in describing the preparation of ceria (p. 332) that rare earth mixtures obtained by the ignition of the mixed oxalates are completely soluble in intir or hydrochloric acid, provided that the ceria does not exceed 45 to 50 per cent of the mixture; pure ceria, however, is insoluble in these acids. The usual explanation of these results is that the cern acts as a feeble acid and combines with the other strong bases present to form salts, these salts are decomposed by a strong acid with the liberation of ceric acid, i.e. ceric hydroxide, which is soluble, as a base, in the excess of strong acid present. If such is the case, the salts must apparently be of the type M<sub>2</sub>O<sub>3</sub> 2CeO<sub>2</sub> or 2M<sub>2</sub>O<sub>3</sub> 3CeO<sub>2</sub> in order to account for the 45-50 per cent limit to the solubility of the cena.

Ceria combines with uranum dioxide. When a dry mixture of cerous and uranyl sulphates is heated with molten magnesium chloride in a covered crucible for fifteen hours, deep blue cubic crystals are produced which can be separated from the accompanying substances by reason of the relative stability

<sup>1</sup> The higher the temperature of formation the higher the density of the crystals. Thus, cenia crystallised from sodium chloride, borax, and potassium sulphate respectively has been found to possess the following densities (Steiba) 7-314, 7-415, and 7-995.

8 K. A. Hofmann and Hoschele, Ber., 1914, 47, 238. For other methods of crystallising ceria, see Nordenskiold, Poog. Annalen, 1861, 114, 612; Didier, An. i. Sci. Ecide norm., 1887, p. 65; Grandeau, Compt. rend., 1885, 100, 1134; Steiba, Compt. rend., 1901, 133,421, 294; Ann. Chim. Phys., 1904, [vin.], 2, 193.

8 Trede and Birithratici, Zeitsch. anny Chem., 1914, 87, 160.

4 Schiffer, Inaugural Dissertation (Munich, 1900); Muthmann and Weiss, Annalen, 1904, 331, 1.

<sup>Schnier, January and January and January and Dissertation (Munich, 1902).
Winkler, Ber., 1891, 24, 873; Holm, Inaugural Dissertation (Munich, 1902).
Marc, Ber., 1902, 35, 2370; R J. Meyer and Koss, Ber., 1902, 35, 3740.
Bekk, Ber., 1913, 46, 2574; Miss Reimer, J. Amer. Chem. Soc., 1915, 37, 1686.</sup> 

towards dilute acids. The composition of the crystals approximates to that required by the compound UO2-2CeO2. A similar deep blue compound may be obtained by precipitating an aqueous solution of uranyl and cerous nitrates with excess of ammonium hydroxide or dilute potassium hydroxide solution;

the precipitate is at first yellow, but soon changes to a denser blue solid.\(^1\) Ceric hydroxide, Cc(OII)\_4—The normal hydroxide is not known. When ammonia or an alkalı hydroxide is added to a solution of a ceric salt, a yellow, gelatinous precipitate of ceric hydroxide is obtained, insoluble in excess of precipitant, it is usually contaminated with basic salt and adsorbed alkali hydroxide. A pure hydroxide may be prepared from ceric aminonium nitrate by precipitation with ammonia from a cold solution, the precipitate being washed, allowed to become nearly dry at a low temperature, and again washed with cold water to remove ammonium nitrate. When dried over potassium hydroxide, its composition corresponds with the formula Ce<sub>2</sub>O(OH)<sub>0</sub> or 2CeO<sub>2</sub> 3H<sub>2</sub>O.<sup>2</sup>

Ceric hydroxide may be conveniently prepared by the oxidation of cerous hydroxide. For this purpose excess of alkali hypochlorite or hypobromite may be added to a cerous salt, or, what amounts to the same thing, precipitation may be effected by alkah hydroxide and the oxidation then accomplished by the use of chlorine or bromine.3 Ceric hydroxide may also be prepared by heating hydrated percence hydroxide to 120° or boiling its aqueous suspension until decomposition of the peroxide has been completed

Ceric hydroxide dissolves in mitric or sulphuric acid with the production of a ceric salt, but reduction to the cerous state usually occurs to a slight extent. According to Brauner, the reduction is complete if the hydroxide has been prepared from perceric hydroxide and its solution in sulphune acid is effected in a platinum dish, but this is denied by Barbieri. 6 Ceric hydroxide

reacts with hydrochloric acid to produce corous chloride, chlorine, and water.

Colloidal veric hydrocide may be prepared by dialysing a 10 per cent. aqueous solution of ceric ammonium nitrate for four or five days. The hydrosol thus obtained is very readily coagulated, when evaporated, it leaves a guinny residue, soluble in hot water. The hydroxide is positively charged.

Perceric hydroxide, CeO<sub>3</sub> ·H<sub>2</sub>O or Ce(OH)<sub>3</sub> O OH —When a cerous salt is treated with a mixture of ammonia and hydrogen perovide, a reddishbrown, gelatinous precipitate of perceric hydroxide is obtained, which has the composition of a hydrated triovide, CeO<sub>3.26</sub>H<sub>2</sub>O.8 According to Pissarjewsky, o the precipitate is produced as follows -

$$Ce(OII)_4 + HO.OII = Ce(OII)_3 O.OII + II_2O.$$

<sup>&</sup>lt;sup>1</sup> K. A. Hofmann and Hoschele, Ber., 1915, 48, 20.

<sup>2</sup> Wyrouboff and Vorneul, Ann Chim Phys., 1906, [vin.], 9, 310, Rannielsberg (Poyg. Annalen, 1859, 108, 40) and Eik (Eritsh file Chem., 1870, [n.], 7, 100) give the same formula, but of Cainelley and Walker, Trans. Chem., Sec. 1888, 53, 50.

<sup>3</sup> Mosander, Phil Mag., 1843, [n.], 23, 211, Popp, Annalen, 1864, 131, 359, Hermann, J. prakt. Chem., 1843, 30, 184, 1864, 92, 113. Stapif, vbul., 1860, 79, 257; Rammelsberg, loc. ed.

loc. crt.
4 Uf. p 391.

 <sup>4 (</sup>f. p. 301.
 Brauner, Zeitsch. anorg. Chem., 1904, 39, 261.
 Barbieri, Atti R. Accad. Linee., 1907, [v.], 16, i. 525.
 W. Biltz, Ber., 1902, 35, 4431; 1904, 37, 1095.
 Hermann, J. prakt. Chem., 1843, 30, 184; Cleve, Bull. Soc. chim., 1885, [ii.], 43, 58; Lecoq de Bousbaudran, Compt. rend., 1885, 100, 605; cf. von Knorie, Zeitsch. angew. Chem., 1897, p. 723; Mengel, Zeitsch. anorg. Chem., 1899, 19, 71, Wyrouboff and Verneuil, Ann. Chim. Phys., 1906, [viii.], 9, 313.
 Pissarjewsky, J. Russ. Phys. Chem. Soc., 1900, 32, 609, Zeitsch. anorg. Chem., 1902, 21, 359.

<sup>31, 359.</sup> 

It is an unstable compound, losing oxygen fairly quickly even at the ordinary temperature. It is rapidly converted into ceric hydroxide when heated with boiling water or when dried at 120° It dissolves in dilute mineral acids, producing a cerous salt, hydrogen peroxide, and oxygen, e.g. —

$$2\text{Ce}(OH)_3 O.OH + 3H_2SO_4 = (e_2(SO_4)_3 + H_2O_2 + O_2 + 6H_2O_3)$$

Ceroceric hydrosulphate.—When cene oxide or hydroxide is heated with fairly concentrated sulphuric acid it is not quantitatively converted into ceric sulphate, oxygen is evolved and part of the cene salt reduced to the cerois state. From the acid solution a beautiful red hexagonal crystalline salt may be readily obtained, and from the mother liquor normal cene sulphate, Ce(SO.), 4H.O. may then be separated in pale vellow crystals.

Ce(SO<sub>4</sub>)<sub>2</sub> 4H<sub>2</sub>O, may then be separated in pale yellow crystals. In addition to the preceding method of preparation, the red crystals may be obtained by mixing cerous sulphate with an excess of ceric sulphate and

crystallising from fairly concentrated sulphuric acid

The hexagonal crystals appear dark orange-coloured when large crystals are viewed in the direction of the principal axis, and orange-yellow when fine needles are similarly examined. Viewed parallel to the basal plane, the crystals appear to have a beautiful red colour. They may be heated to 230° without losing anything but their water of crystallisation.

The composition of these hexagonal crystals has been a matter of dispute for many years, the formula that have been assigned to the substance are

given in the appended table -

| _ <del> </del>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |                                                                                                                                                                                                                                                                                                                                                                     | ,                                                                                                    |
|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------|
| Formula                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | Anthouty                                                                                                                                                                                                                                                                                                                                                            | Date                                                                                                 |
| $\begin{array}{c} Ce_1O_0 \text{ $SO_2.27H,0} \\ SCe(SO_4) \text{ $\ThetaC_2/SO_4\}_2 \text{ $27H,0}} \\ SCe(SO_4) \text{ $\ThetaC_2/SO_4\}_2 \text{ $27H,0}} \\ SCe(SO_4)_2 \text{ $Ce_2/SO_4\}_2 \text{ $27H,0}} \\ SCe(SO_4)_2 \text{ $Ce_2/SO_4\}_1 \text{ $21I_2SO_4$_25H,0}} \\ 2Ce(SO_4)_2 \text{ $14Ce_2/SO_4\}_1 \text{ $11I_2SO_4$_25H,0}} \\ SCe(SO_4)_2 \text{ $Ce_2/SO_4\}_2 \text{ $21H,0}} \\ SCe(SO_4)_2 \text{ $Ce_2/SO_4\}_3 \text{ $21H,0}} \\ SCe(SO_4)_2 \text{ $Ce_2/SO_4\}_3 \text{ $21H_2O$}} \\ 2Ce(SO_4)_2 \text{ $Ce_2/SO_4\}_3 \text{ $21H_2O$}} \\ 2Ce(SO_4)_2 \text{ $Ce_2/SO_4\}_3 \text{ $21H_2O$}} \\ 2Ce(SO_4)_2 \text{ $Ce_2/SO_4\}_3 \text{ $11H_2O$_4$_26H_2O$}} \\ 2Ce(SO_4)_2 \text{ $Ce_2/SO_4\}_3 \text{ $11H_2O$_4$_26H_2O$}} \\ 2Ce(SO_4)_2 \text{ $Ce_2/SO_4\}_3 \text{ $11H_2O$_4$_26H_4O$}} \\ SCe(SO_4)_2 \text{ $Ce_2/SO_4\}_3 \text{ $11H_2O$_4$_21H_4O$}} \\ SCe(SO_4)_2 \text{ $Ce_2/SO_4\}_3  $11H_2O$_4$_21H_4O$_4$_4$_4$_4$_4$_4$_4$_4$_4$_4$_4$_4$_4$$ | Hermann <sup>1</sup> Rannnelsberg <sup>2</sup> Czudnowicz <sup>3</sup> Hermann <sup>4</sup> Zschiescho <sup>5</sup> Rrunnersberg <sup>6</sup> Mendelceff <sup>†</sup> John <sup>8</sup> Brauner <sup>10</sup> Brauner <sup>10</sup> Muthanann and Stutzel <sup>11</sup> Meyer and Aufrecht <sup>12</sup> Brauner <sup>17</sup> Wyrouboff and Verneurl <sup>14</sup> | 1843<br>1859<br>1860<br>1864<br>1869<br>1873<br>1873<br>1874<br>1895<br>1895<br>1900<br>1904<br>1904 |
|                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |                                                                                                                                                                                                                                                                                                                                                                     |                                                                                                      |

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1 Hermann, J. makt. Chem., 1843, 30, 184
2 Rammelsberg, Popp. Annalen, 1859, 108, 40.
3 Czudnowicz, J. makt. Chem., 1860, 80, 16.
4 Hermann, J. makt. Chem., 1864, 92, 113
5 Zschresche, J. prakt. Chem., 1869, 107, 65
6 Rammelsberg, Ber., 1873, 6, 568, Annalen, 1873, 168, 45
7 Mendeléeff, Ber., 1873, 6, 568, Annalen, 1873, 168, 45
8 John, Bull. Soc. chem., 1894, [11], 21, 533
9 Brauner, Chem. New., 1895, 71, 283
10 Rhauner, Bull. Intern. Anal. Sci. de l'Empereur François Joseph I., 1895.
11 Muthmann and Stutzel, Ber., 1904, 33, 1763
12 R. J. Meyer and Antiecks, Ber., 1904, 37, 140
13 Brauner, Zeitsch. anorg. Chem., 1904, 39, 2012
14 Wyrouboff and Verneuil, Ann. Chem. Phys., 1806, [viii.], 9, 293
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The substance contains both cerous and ceric sulphates as well as sulphuric - acid, and the latter cannot be eliminated at 130° in vacuo. It may therefore be described as a ceroceric hydrosulphate. The cerous sulphate may be replaced by the sulphates of the other rare earth elements, and Brauner has prepared the lanthanum, praseodymium, and neodymium salts. They are hexagonal, and isomorphous with one another and with the cerous salt According to Brauner, they are acid salts of a complex cerisulphuric acid, H<sub>4</sub> Ce<sup>V</sup>(SO<sub>4</sub>)<sub>4</sub>. 12H<sub>2</sub>O, of the type HM<sup>III</sup>.Co<sup>IV</sup>(SO<sub>4</sub>)<sub>4</sub>. 12H<sub>2</sub>O (M<sup>III</sup> = Ce, I.a, Pr, or Nd). 1

It will be noticed that the formula assigned to coroceric hydrosulphate by Meyer and Aufrecht only differs from Brauner's formula (1904) in the amount of water of crystallisation present. Too much significance should not, however, be attached to this agreement, for the analytical data upon which the formulæ are based are of slender value, and at the present time the composition and constitution of this interesting compound must be regarded as undetermined.2

Ceric sulphate, Ce(SO<sub>4</sub>)<sub>2</sub>—Ceria is quantitatively converted into ceric sulphate when heated on a sand-bath with concentrated sulphuric acid. The salt, which is insoluble in the excess of acid, may be washed with glacial acetic acid and dried over potash, when it is obtained as a deep yellow, crystalline powder When heated in the air, slight loss of weight is observed

| Per cent. of                                                   | Calculated from the Formula of  |                                 |                                 | Found by               |                                 |                                 |
|----------------------------------------------------------------|---------------------------------|---------------------------------|---------------------------------|------------------------|---------------------------------|---------------------------------|
| Ter cont. or                                                   | Brauner                         | W. and V.                       | H.F.V.L                         | Brau                   | inei                            | W. and V.                       |
| Ce <sub>2</sub> O <sub>3</sub> Active oxygen . SO <sub>3</sub> | 87 25<br>0:91<br>36:31<br>25:53 | 87 75<br>0.79<br>35.63<br>25 88 | 87·27<br>0 78<br>36·29<br>25·66 | 36·57<br>0·82<br>26·08 | 36 15<br>0 84<br>36 78<br>26 13 | 37·57<br>0·79<br>86·16<br>25·51 |

which serves to illustrate the difficulties attaching to the investigation of the composition

 $<sup>^1</sup>$  It may be remarked that the acid sulphate of thorium has the composition  $Th(SO_4)_9, H_2SO_4$  or  $H_2Th(SO_4)_5,$  and not  $H_4Th(SO_4)_6.$  On the other hand, the acid sulphates of the rare earth elements have formulae of the type  $M_4^{II}(SO_4), 3H_2SO_4$  or  $H_4[M_1^{II}(SO_4)_5]$ , the ceric salt of which would be Cely MIII(SO<sub>4)214</sub> or 3CelV(SO<sub>4)2</sub> 2MIII(SO<sub>4)3</sub>, in harmony with the formula of Wyrouboff and Verneull; it is then difficult, however, to account for the

the formula of Wyrouboff and Verneuil; it is then difficult, however, to account for the extra sulphunic acid present in the molecule.

If cerocenic hydrosulphate is a "complex" salt, it is not a very stable complex, for the addition of alkali leads to the initial precipitation of the violet cerocenic hydroxide

Thus, oven the atomic latio Cel<sup>17</sup> Cel<sup>11</sup> is uncertain. The analytical data given by Myrouboff and Verneuil agree very well with the result Cel<sup>17</sup> Cel<sup>11</sup> 3:4, and the data given by Moyer and Aufrecht and by Brauner agree at least as well with this ratio as with the ratio 1 1, which they adopt. Brauner's analyses of the lanthanum, praceodymium, and neodymium salts are also unsatisfactory, as Wyrouboff and Verneuil have justly remarked.

It is, perhaps, worth while to point out that the formula 3Ce(SO<sub>1</sub>), 2Ce<sub>2</sub>(SO<sub>2</sub>), 2Ce<sub>3</sub>(SO<sub>4</sub>), 2H<sub>3</sub>SO<sub>4</sub>.

42H<sub>4</sub>O agrees with the analytical data given by Brauner and by Wyrouboff and Verneuil at least as well as do then own formulæ. This is readily seen from the following table of results:—

of occorn hydrosulphate.

Finally, it may be mentioned that Brauner has described a ceroceric sulphate, Collifical (SO<sub>4</sub>)<sub>1</sub>, 44H<sub>2</sub>O, and the corresponding lanthanoceric salt

Rammelsberg, Ber., 1873, 6, 84, Mendeléeff, Annalen, 1873, 168, 45; Ber., 1873, 6, . 558; Muthmann and Stutzel, Ber., 1900, 33, 1708; R. J. Meyer and Aufrecht, Ber., 1904, 37, 140; Brauner, Zeitsch. anorr. Chem., 1904, 39, 261; Wyrouboff and Veneul, Ann. Chim. Phys., 1906, [viu.], 9, 311; Barbieri, Atti R. Accad. Lincei, 1907, [v.], 16, i. 525.

at 155°, and when heated to constant weight at 195° the basic sulphate 3CeO<sub>2</sub>.4SO<sub>3</sub> is produced On raising the temperature, further loss of sulphur trioxide occurs at 250°, and at 300° the loss in weight is continuous until cerium dioxide is left.1

Ceric sulphate is very soluble in water, in which it forms a yellowishbrown, unstable solution. A concentrated solution of the salt in dilute sulphuric acid may be prepared either by dissolving the anhydrous salt in dilute sulphuric acid or by carefully dissolving cenic hydroxide in the same medium. When concentrated over sulphuric acid, sulphur-yellow, orthorhombic 2 crystals of the tetrahydrate, Ce(SO<sub>1</sub>)<sub>2</sub> 4H<sub>2</sub>O, are obtained (bipyramidal;  $\ddot{a} \cdot b : c = 0.717 \cdot 1 = 0.171$ )

Ceric sulphate in solution readily hydrolyses, and, according to Wyrouboff and Verneuil, when diluted largely or warmed, a crystalline basic salt of the composition 2CeO, 2SO, 5H2O is obtained, 4 from a dilute ico-cold solution the compound  $4\text{CeO}_2$   $3\text{SO}_3$   $12\text{H}_2\text{O}$  has been prepared, and from a saturated solution at 0° C the compound  $2\text{CeO}_2$   $3\text{SO}_3$   $4\text{H}_2\text{O}^{-6}$  A physicochemical study of the hydrolysis has been made by Spencer, who could only prepare and characterise one basic salt at 25° C., viz CeO<sub>2</sub> SO<sub>1</sub> 2H<sub>2</sub>O  $^6$ 

Cover potassium sulphate,  $Ce(SO_1)$ ,  $2K_aSO_1$ ,  $2H_aO_5$  is obtained in orange yellow, monoclinic crystals (a:b:c=1.216-1.2.093),  $\beta=100^{\circ}$  40'), searcely soluble in water, by adding a solution of potassium sulphate to an acid solution of coric sulphate 8

Ceric ammonium sulphate, Ce(SO<sub>1</sub>)<sub>2</sub>,2(NII<sub>1</sub>)<sub>2</sub>SO<sub>1</sub>,2II,O, separates in yellow crystals when an acid solution of the mixed sulphates is evaporated; it is followed by orange-red crystals of the monochine (a  $b \cdot c = 0.6638.1 - 0.7838$ ;

 $\beta = 96^{\circ} 44')^{\circ} \text{ double salt Ce(SO_1)}_2 3(\text{NH}_1)_2 \text{SO}_1 2 \text{H}_2 \text{O}^{-10}$ Ceric silver sulphate, 5Ce(SO\_1)}\_2 3Ag\_SO\_1 is obtained as an orange yellow, crystalline precipitate by adding silver intrate to a strongly acid solution of The salt slowly combines with 2H2O, and is decomposed by ceric sulphate boiling water.11

Ceric selenite, Ce(SeO<sub>3</sub>)<sub>2</sub>, may be prepared by heating cerous intrate and selenious acid with boiling concentrated mitrie acid. It is an orangeyellow powder, insoluble in water but slightly soluble in intiic acid 13

Ceric chromate, Ce(CrO<sub>1</sub>), 2H<sub>2</sub>O, is obtained by heating ceric hydroxide with aqueous chromic acid. It is a bright scarlet crystalline salt that cannot be heated above 180° without decomposing - It is decomposed by water, losing chromic acid and turning orange in colour 13

<sup>1</sup> J. F. Spencer, Trans. Chem. Sur., 1915, 107, 1265.
2 Slavik, Zettsch. anorg. Chem., 1904, 39, 280

1 According to Wyrouboff and Verneurl (Ann. Chem. Phys., 1906 [viii] 9, 312), the formula is 2Cc(SO<sub>2</sub>), 7H<sub>2</sub>O.

4 Wyrouboff and Verneurl. Ann. Chem. Phys., 1908, [viii], 9, 312.

5 Hauser and Wirth, Zettsch. anorg. Chem., 1908, [viii], 9, 312.

6 J. F. Spencer, Trans. Chem. Sur., 1915, 107, 1265.

7 Marignac, Ann. Min., 1859, [v.]. 15, 221

8 Scheerer, Pogg. Annalen., 1812, 56, 482, Hermann, J. puckt. Chem., 1813, 30, 184; Rammelsberg, Pogg. Annalen., 1859, 108, 40, Marignac, loc. cit.

9 (icipel, Zettsch. Kryst. Min., 1902, 35, 608.
10 Rammelsberg, Pogg. Annalen., 1859, 108, 10, Bcr., 1873, 6, 84, Mondeléeff, Annalen., 1873, 168, 45; Geipel, loc. cit.

11 Pozzi-Escot, Compt., rend., 1913, 156, 1074.

12 Barbiert and Calzolari, Ber., 1910, 43, 2214.

13 Browning and Flora, Amer., J. Sei., 1903, [iv.], 15, 177, cf. Bricout, Compt. rend., 1894, 118, 145; Bohm., Zeitsch. angew. Chem., 1902, 15, 372, 1282.

Ceric molybdate.—A number of salts of a cerimolybdic acid have been prepared, analogous to the complex molybdates of zirconium and thorum. Ammonium cerimalyhdate, (NH<sub>1)x</sub>[Cc(Mo<sub>2</sub>O<sub>2</sub>)<sub>6</sub>] 8H<sub>2</sub>O, is a yellow crystalline substance, yielding yellow solutions which are stable towards hydrogen peroxide and give a precipitate with oxalic acid only after prolonged boiling. The acid ammonium salt (NH<sub>4</sub>)<sub>6</sub>H<sub>2</sub>[Ce(Mo<sub>2</sub>O<sub>7</sub>)<sub>6</sub>] 10H<sub>2</sub>O, the silver salt, Ag<sub>8</sub>[Ce(Mo<sub>2</sub>O<sub>7</sub>)<sub>6</sub>], and the acul aniline salt are also known.

# CERIUM AND THE NITROGEN GROUP.

Ceric nitrate, Ce(NO<sub>3</sub>), has not been isolated. By evaporating a solution of ceric hydroxide in concentrated nitric acid, in the presence of a considerable quantity of calcium intrate, red monoclinic crystals (a:b:c= $1.7834 \cdot 1 \quad 1.0465$ ,  $\beta = 90^{\circ} 48'$ ) of basic cert intrate,  $20'e(OH)(NO_3)$ ,  $9H_2O$ , can be obtained. The aqueous solution of the salt readily undergoes hydrolysis (p. 332).3

Ceric nitrate forms an isomorphous series of double nitrates with the nitrates of potassium, rubidium, casium, and ammonium They are of the type Ce(NO<sub>3</sub>), 2MNO<sub>3</sub>, and form anhydrous, 1 orange-red, monoclinic crystals:-

The salts are readily soluble in water and alcohol, but much less soluble in fairly concentrated intric acid. In aqueous solution they are hydrolysed in a similar manner to cere initiate. It is probable that the solutions contain not only the ions Ce<sup>...</sup>, M\*, and NO<sub>3</sub>', but also the complex ion Ce(NO<sub>3</sub>)<sub>6</sub>"; indeed, in alcoholic solution it has been shown by migration experiments that a coloured cation is present.

Ceric ammonium nitrate, Ce(NO<sub>3</sub>)<sub>1</sub> 2(NH<sub>1</sub>)NO<sub>30</sub> is of considerable practical importance for the preparation of pure ceria (see p. 336). Indeed, from the practical point of view, it is the most important of the ceric salts. Its solubility in water is as follows (in grams of salt per 100 grams of solution):-

Temp ° C. 
$$25^{\circ}$$
  $35 \cdot 2^{\circ}$   $45 \cdot 3^{\circ}$   $64 \cdot 5^{\circ}$   $85 \cdot 6^{\circ}$  Grams of  $Ce(NO_3)_1 \cdot 2(NH_1)NO_4 \cdot 58 \cdot 5 \cdot 62 \cdot 8 \cdot 64 \cdot 5 \cdot 66 \cdot 8 \cdot 69 \cdot 4$ 

At temperatures above 60° partial reduction to cerous nitrate takes place in the aqueous solution.

It may readily be prepared by the electrolytic oxidation of cerous ammonium nitrate in a divided cell 8

<sup>1</sup> Barbiell. Atti R. Accad. Lines, 1914, [v.], 23, 1-805.
2 von Lang and Haitinger, Annalen, 1907, 351, 450.
3 R. J. Meyer and Jacoby, Ber., 1900, 33, 2135. Zeitsch. anory Chem., 1901, 27, 359; these authors give the water of crystallisation as 3H<sub>2</sub>O and not 4½H<sub>2</sub>O.
3 Meyer and Jacoby, loc cit.; Wolff, Zeitsch anory Chem., 1905, 45, 89, cf. Holamann, J. pratt. Chem., 1858, 75, 321; Muthmann and Rolig, Zeitsch. anorg. Chem., 1898, 45, 450.

<sup>16, 450</sup>Sachs, Zeilsch. Kryst. Min., 1901, 34, 162
Sachs, Zeilsch. Kryst. Min., 1901, 34, 162
Meyer and Jacoby, loc. cit. See also Barbieri, Atti R. Accad. Lincei, 1907, [v.], 16, i. 395. Wolff, loc. cit.

Plancher and Barbiers, Atti'R Accord. Lincei, 1904, [v.], 14, i. 119.

# THE RARE EARTH ELEMENTS.

'Ceric nitrate also forms an isomorphous series of double salts with the nitrates of the bivalent metals magnesium, zinc, nickel, cobalt, and manganese; they are of the type  $Ce(NO_8)_4.M(NO_8)_2.8H_2O$ , and must be crystallised from concentrated nitric acid. The crystals are monoclinic and isomorphous with the corresponding double thorium nitrates :- 2

Ceric orthophosphate. - When sodium phosphate is added to a slightly acid solution of a ceric salt, a yellow precipitate is obtained. According to Hartley, the composition of this piccipitate (dried in vacuo over sulphuric acid) is in accordance with the formula 2Co<sup>1</sup>.H(PO<sub>1</sub>), 25H<sub>2</sub>O, but this cannot be considered as definitely established.3 Cente pyrophosphate, Cel'2O7.xH2O(1), like the corresponding thorium salt, is practically insoluble in dilute mineral acids. It is obtained as a yellow precipitate by warming a solution of cerous pyrophosphate in hydrochloric acid with bromine.4

Ceric dihydrogen arsenate, Ce(II,AsO<sub>1</sub>), 4H<sub>2</sub>O<sub>2</sub>, may be prepared by heating cerous nitrate (1 mol) and arsenic acid (4 mols.) with concentrated nitric acid for several hours, distilling off most of the nitric acid, and allowing the liquid to crystallise. The salt separates in white needles. When it is dissolved in the minimum quantity of concentrated intric acid and the cold solution diluted with water, a white, crystalline precipitate of ceric monohydrogen arsenate, Ce(IIAsO<sub>1</sub>)<sub>2</sub>.6H<sub>2</sub>O, is obtained <sup>5</sup>

# CERIUM AND THE CARBON GROUP.

Ceris carbonate -A yellow, gelatinous precipitate, presumably of a basic ceric carbonate, is obtained by double decomposition between a coric salt and an alkali carbonate. The precipitate dissolves is addy in concentrated potassium carbonate solution, forming a yellow solution that is stable in the air but which may be oxidised with hydrogen peroxide.6 The basic cerio carbonates, 2CeO, 3CO, and 2CeO, CO, are obtained (combined with potassium carbonate)? by heating perceric potassium carbonate to 240° and 360° respectively.7

Perceric carbonate.—Corous carbonate dissolves to a considerable extent in concentrated potassium carbonate solution. When hydrogen peroxide is added to the solution, the latter turns blood-red in colour owing to oxidation, and the colour darkens with the addition of hydrogen peroxide up to a certain point; beyond this, the addition of more perovide lightens the colour and throws down an orange-yellow precipitate, and by the addition of sufficient reagent all the cerium is precipitated.

<sup>1</sup> Bunsen and Jegel, Annalen, 1858, 105, 40, 45; Holymann, loc. cit.; Meyer and

Bunsen and Jegel, Annaten, 1888, 105, 40, 45; Hollmann, toe. cu.; steper and Jacoby, loc. cit.

Geipel, Zeitsch. Kryst. Min, 1902, 35, 608.

Hartley, Trans. Chem. Soc., 1882, 41, 202.

Carney and Campbell, J. Amer. Chem. Soc., 1914, 36, 1131, Rosenheim and Triantaphyllides, Ber., 1915, 48, 582.

Barbieri and Calzolari, Ber., 1910, 43, 2214.

Job, Ann. Chim. Phys., 1900, [vii.], 20, 205

Job, loc. cit.

It is much less soluble in sodium or ammonium carbonate.

When the colour reaches its maximum intensity, all the cerum is present in solution as perceric carbonate The solution of perceric carbonate in potassium carbonate may be prepared from ceric carbonate as well as from cerous carbonate, by oxidation with hydrogen peroxide. By a suitable process of orystallisation, Job obtained perceric potassium carbonate, Ceg(COg), Og.

4K<sub>2</sub>CO<sub>3</sub>.12H<sub>2</sub>O, from this solution in dark-red, triclinic crystals.

This curious double salt is remarkably stable—It may be dehydrated at 110°, but withstands a temperature of 200° for several hours without decomposition being perceptible. At 240°, however, it loses oxygen, leaving

a residue of a basic ceric carbonate.1

By modifying Job's method of preparation in certain particulars, Meloche 2 has prepared a crystalline percene potassium carbonate of the composition  $\text{Ce}_2\text{O}_4(\text{CO}_3)_2$   $4\text{K}_2\text{CO}_3$   $12\text{H}_2\text{O}$ . This compound readily loses part of its water when exposed to dry air, and may be almost completely dehydrated at 110°-120° without loss of available oxygen.

As already mentioned, the addition of hydrogen perovide to perceric potassium carbonate solution causes all the cerium to be thrown out of solution as an orange-yellow precipitate. This precipitate appears to be a derivative of CeO, and is very unstable. When covered with a concentrated solution of potassium carbonate it slowly evolves oxygen, and beautiful red crystals of a percenc potassium carbonate are formed. According to Job, this is the best method for preparing the double salt, according to Baur, however, the crystals formed are rather different in composition from those described by Job, and have the formula Ce<sub>2</sub>O<sub>1</sub>(CO<sub>3</sub>), 4K,CO<sub>3</sub> 10H<sub>2</sub>O.<sup>3</sup>

In alkaline solution the perceric potassium carbonates have three atoms. of available oxygen per two atoms of cerium. In acid solution, however, only one-third of this oxygen is available, the remainder being set free in the gascons state, e.g -

$$\begin{aligned} \text{Ce}_2\text{O}_1(\text{CO}_3)_2 & 4\text{K}_2\text{CO}_3.12\text{H}_2\text{O} + 7\text{H}_2\text{SO}_4 \\ &= \text{Ce}_2(\text{SO}_4)_3 + \text{O}_2 + \text{H}_2\text{O}_2 + 4\text{K}_2\text{SO}_1 + 6\text{CO}_2 + 18\text{H}_2\text{O}_2 \end{aligned}$$

A solution of ceric carbonate in potassium carbonate is quite stable towards air or oxygen. On the other hand, the corresponding cerous solution readily absorbs oxygen, passing into perceric and not into ceric carbonate. Simultaneously with this reaction another change occurs, namely, the interaction of the percenc carbonate produced with the unchanged cerous carbonate to produce ceric carbonate; and by varying the conditions of experiment the relative speeds of those two reactions may, within certain limits, be altered at will. This auto-oxidation of cerous carbonate may be used to affect the oxidation of various substances by the air, a small quantity of cerous salt acting as the oxygen-carrier. It is only necessary for this purpose that the "acceptor" shall be able to reduce both ceric and perceric carbonates to the cerous state; glucose is one such substance. When, however, the "acceptor" can only reduce perceric carbonate to the ceric state, the cerous salt soon becomes quantitatively transformed into ceric salt, when the auto-oxidation ceases According to Engler, the auto-oxidation of cerous tarbonate takes place in two stages, as follows:-

$$\begin{array}{ll} \text{(i.)} & \operatorname{Ce_2(CO_3)_3} + 2\operatorname{H_2O} + \operatorname{O_2} & = \operatorname{Ce_2(CO_3)_3(OH)_2} + \operatorname{H_2O_2} \\ \text{(ii.)} & \operatorname{Ce_2(CO_3)_3(OH)_2} + 2\operatorname{H_2O_2} = \operatorname{Ce_2(CO_3)_3O_3} + 3\operatorname{H_2O} \end{array}$$

Job, Ann. Chim. Phys., 1900, [vii\*], 20, 205.
 Meloche, J. imer. Chem. Soc., 1915, 37, 2338, 2645.
 Baui, Zeilsch. anorg. Chem., 1902, 30, 256.

the initial products of oxidation being basic ceric carbonate and hydrogen peroxide, which then interact to produce the percoric compound.

Ceric acetate. - The normal salt is not known. A weakly acid solution of a ceric salt is completely precipitated as yellow basic ceric acetate when heated to boiling with excess of an alkali acetate. An aqueous solution of cerous acetate is said to be ovidised to cone acetate by ozone.2

Basic perceric acetate is obtained as an orange-brown precipitate when hydrogen peroxide and an alkali acetate are added to a solution of a cerous salt The precipitate thus obtained is apparently a derivative of a peroxide (Cc(OII)2.O.II) analogous to the peroxides of the other rare earth elements, since the atomic ratio of cerium to active oxygen (i.e. oxygen in excess of that required for a cerous salt) is Ce. O 3 The formation of basic perceric acetate may be used as a test for cerruin or as a means of separating cerium from the other care earth elements 4 The precipitate is converted into basic ceric acetate when dried at 120'.

Ceric oxalate, ('e(C<sub>2</sub>O<sub>1</sub>), 7H<sub>2</sub>O (?) — ('eric oxalate is obtained as an orange-yellow gelatmous precipitate when cold, aqueous solutions of ceric ammonium intrate and ammonium oxalate are mixed. The precipitate is difficult to filter and wash, and when attempts are made to dry it on a porous plate, considerable decomposition occurs, apparently a mixture of cerous oxalate, Ce<sub>2</sub>(C<sub>2</sub>O<sub>4</sub>), 10H<sub>2</sub>O, and ceric oxalate, Ce(C<sub>2</sub>O<sub>4</sub>), 7H<sub>2</sub>O, is produced, from which ceric oxalate may be extracted with aqueous ammonium oxalate.5

Cerous oxalate is a very unstable salt, readily losing carbon dioxide and becoming converted into cerous oxalate:-

$$2\mathrm{Ce}(\mathrm{C_2O_1})_2-\mathrm{Ce_2}(\mathrm{C_2O_1})_3+\mathrm{CO_2}.$$

It is readily soluble in ammonium oxalate solution, thereby resembling thorium ovalate The orange-yellow solution deposits cerons oxalate on standing, the decomposition being rapid when the solution is warmed. The cerium may be rapidly and quantitatively precipitated from the solution by the addition of sulphurous acid 6

Ceric acetylacetonate, Ce(CH, CO CH, CO, CH, ), may be prepared by shaking a suspension of ceric hydroxide in water with excess of acetylacetone for several days. Deep red needles of the hydrated compound (1111,0) are thus formed They are dried in vacuo and crystallised from carbon tetrachloride, when black, lustious crystals of the anhydrous substance separate out, m.p. 171–172'. The acetylacetonate is slightly soluble in water, which hydrolyses it slightly, hydrolysis may be repressed by the addition of a little acetylacetone. The compound dissolves in most organic media, forming deep red solutions that are readily reduced.7

7. .

For further information on the auto-oxidation of cerous salts, see Job, Compt. rend., 1898, 126, 246; 1899, 128, 178, 1098, 1902, 134, 1052, 1903, 135, 45. Ann. Chim. Phys., 1900, [vii.], 20, 205, Baur, Zeitsch. anorg. Chem., 1902, 30, 251. Ibr., 1903, 36, 3038, 1904, 37, 795; Engler and Wohler, Zeitsch. anorg. Chem., 1902, 20, 1, Engler, Ber., 1903, 36, 2842, 1904, 37, 49, 3268. Engler and Wersberg, Kritische Stadten über die Vorgange der Auto orydation (Vieweg, Brunswick, 1904).
 Job, Compt. rend., 1903, 136, 45.
 Wyrouboff and Veineuil, Ann. Chim. Phys., 1906, [viii], 9, 314, cf. Meloche, loc. cit.
 Popp, Annalen, 1864, 131, 359; R. J. Meyer and koss, Ber., 1902, 35, 672.
 Brady and Little; unpublished experiments.
 See Orloff, Chem. Zeit., 1906, 30, 733; 1907, 31, 562.
 Job and Goissedet, Compt. rend., 1913, 157, 50.

# ALUMINIUM AND ITS CONGENERS.

### CERIUM AND BORON.

Ceric borate, CeO<sub>2</sub> B<sub>2</sub>O<sub>3</sub>, may be prepared, according to Holm, by heating ceria with an excess of boric anhydride in a petroleum injectorfurnace and extracting the mass with water; it is described as a yellow powder, insoluble in water and dilute mmeral acids.1 Guertler,2 however, states that this method of preparation only yields cerous metaborate, Ce(BO2)8.

### LANTHANUM.

Symbol, La. Atomic weight, 139.0 (0=16).

Lanthanum is one of the most abundant of the rare earth elements. Its occurrence, history, preparation, atomic weight, and homogeneity have been

already discussed in Chapters X. and XI.

Lanthanum 3 is a tin-white metal of density 6.155. It is malleable, but not ductile Its specific and atomic heats (0°-100°) are 0 0449 and 6.23 respectively; 4 its melting-point is 810°. Measurements made with material of doubtful purity indicate that lanthanum is feebly paramagnetic,5 but this is questionable.6

Lanthanum tarnishes rapidly, even in dry air. It burns in air at 440°-460°. The chemical properties of the metal, so far as they are known, are

very similar to those of cerium.

ŗ

Alloys.-Little is known of the alloys of lanthanum. Lanthanum alloys with aluminium, and if excess of the latter is taken, and subsequently removed with sodium hydroxide, the compound LaAl, is obtained. Its density is 3.923 and its heat of formation 97.8 Cals.; it resembles the corresponding cerium compound.7

### COMPOUNDS OF LANTHANUM.

The salts of lanthanum are derived from the colourless basic oxide La<sub>2</sub>O<sub>8</sub>, . and, if derived from colourless acids, are themselves colourless Their aqueous solutions are devoid of absorption spectra

The following values for the equivalent conductivities  $\lambda$  of a number of lanthanum salts at 25° are in harmony with the view that the salts are derived from a fairly strong traced base (v = dilution in litres per gramequivalent):—8

<sup>1</sup> Holm, Inaugural Dissertation (Munich, 1902).
2 Guertler, Zeitsch. anorg. Chem., 1904, 40, 225.
3 See pp. 229-220. Hillebrand and Norton, Pogy. Annalca, 1875, 155, 633, 156, 466; Muthmann and Kraft, Annalca, 1902, 325, 261; Muthmann and Weiss, ibid., 1904, 331, 1 Muthmann and Scheidmandel, ibid., 1907, 355, 116.
4 Hillebrand, Pogg. Annalca, 1876, 158, 71.
5 Owen, Ann. Physik, 1912, [iv], 37, 657; Proc. K. Akad. Wetensch. Amsterdam, 1911, 14, 637.
7 Pure lanthana is damagnetic (p. 257).
7 Muthmann and Bock, Annalca, 1904, 331, 46.
7 Anfrecht, Inaugural Dissertation (Berlin, 1904), Muthmann, Ber., 1898, 31, 1829.
7 Zeitsch, physikal. Chem., 1899, 30, 193; Holmberg, Arkiv Kem. Min. Geol., 1903, 1, 2, A. A. Noyes and J. Johnston, J. Amer. Chem. Soc., 1909, 31, 987; A. Heydweiller, Physikal. Chem., 1915, 89, 281.

# THE RARE EARTH ELEMENT

| 2 : 2                             |             |       |       |        |               |               |         |        |
|-----------------------------------|-------------|-------|-------|--------|---------------|---------------|---------|--------|
| LaCl                              | ข 🖚         | 21.3  | 42.6  | 85.2   | 170.6         | 341.1         | 682.2   | 1364.5 |
|                                   | $\lambda =$ | 101.7 | 1086  | 114.9  | $122 \cdot 3$ | 126.5         | 131.9   | 136.4  |
| La(NO <sub>3</sub> ) <sub>3</sub> | v=          | 32    | 64    | 128    | 256           | 512           | 1024    |        |
| , ( 3/3                           | λ=          | 98.6  | 105 4 | 112.8  | 1181          | $124 \cdot 1$ | 126.5   | •      |
| $La_2(SO_4)_3$                    | . v =       |       | 67.72 | 135.44 | 270 88        | 541.76        | 1083 52 |        |
| 4\4/.                             | λ=          | 43 58 | 50.30 | 58.49  | 68.89         | 80.93         | 96.26   | •      |

Noyes and Johnston have measured the equivalent conductivities of dilute solutions of lanthanum nitrate and sulphate over a wide range of temperature, and Johnston 1 has calculated from the results that the ionic conductivity of the lanthanum ion varies with the temperature as follows -

At 25° the percentage dissociation of the nitrate and sulphate is as follows (Noves and Johnston):--

| $v$ $\Gamma000\eta$ . | <b>x</b><br>0 | $\begin{array}{c} 500\ 0 \\ 0.002 \end{array}$ | 80 00<br>0 01:25 | $\begin{array}{c} 20.00 \\ 0.050 \end{array}$ | 10·00<br>0·100 | 5 00<br>0 200 |
|-----------------------|---------------|------------------------------------------------|------------------|-----------------------------------------------|----------------|---------------|
| $La(NO_3)_3$          | 100           | 90·4                                           | 80·4             | 70 0                                          | 64·4           | 58 6          |
| $La_2(SO_1)_3$        | 100           | 15·0                                           | 27 8             | 19 0                                          | 15 8           | 13 2          |

Numerous compounds of lanthanum have been prepared and described by Cleve,2 and by Frenchs and Smith.8

Thermochemistry of Lanthanum -The following results have been

Lanthanum hydride, LaH<sub>8</sub>, has been described (p. 251).

# LANTHANUM AND THE FLUORINE GROUP.

Lanthanum fluoride, LaF<sub>3</sub>. — See p. 252. The hemihydrate, 2LaF<sub>3</sub>. H<sub>2</sub>O, and the acid fluoride, 2LaF<sub>3</sub>. 3HF, have been described (Cleve; Frerichs and Smith).

Lanthanum chloride, LaCl3. - For methods of preparation and

<sup>71, 484.</sup>Matignon, Ann. Chim. Phys., 1906, [viii.], 8, 426 & Bourion, ibid., 1910, [viii.], 21,

properties of the anhydrous salt, see p. 252. It is a white, crystalline, hygroscopic solid, readily soluble in water.

From an aqueous solution of the chloride, large triclinic crystals (a:b:c=1:1593.1:0:8659; a=91° 3',  $\beta=114°$  28',  $\gamma=88°$  12') of the heptahydrate, LaCl<sub>3</sub> 7H<sub>2</sub>O, may be obtained <sup>1</sup> The double salts listed below have been prepared, in addition to those containing metals of Groups IV. and V. .-

> LaCl<sub>3</sub>,3CsCl.4H<sub>2</sub>O(15H<sub>2</sub>O) 2LaCl<sub>3</sub>,9HgCl<sub>3</sub>,24H<sub>2</sub>O<sup>2</sup> LaCl<sub>3</sub> 3Hg(CN)<sub>2</sub> 8H<sub>2</sub>O 3

 $\begin{array}{c} {\rm LaCl_{*}\,AuCl_{*}\,10H_{2}O^{4}} \\ {\rm 2LaCl_{*}\,3AuCl_{*}\,21H_{2}O(?)^{5}} \end{array}$ 21aCl<sub>3</sub>3PtCl<sub>2</sub>18H<sub>2</sub>06 IaCl<sub>3</sub>PtCl<sub>1</sub>13H<sub>2</sub>04 21aCl<sub>3</sub>3PtCl<sub>1</sub>24H<sub>2</sub>05

Lanthanum oxychloride, LaOCI, has been described 7

Lanthanum bromide, LaBr<sub>1</sub> -- The anhydrous salt has been prepared (p. 255) and also the heptahydrate, LaBr, 7H,O (Cleve), the auxibromide, LaBr, AuBr, 9H, O (Cleve), and lanthanum zine bromide, 2LaBr, 3ZnBr, 39H, O(1) (Frerichs and Smith).

Lanthanum iodide, Lal<sub>3</sub>, is not known, but lanthanum zinc rodide, 2Lal<sub>3</sub>,3Znl<sub>3</sub>,27H<sub>2</sub>O, has been described (Frenchs and Smith)

Lanthanum perchlorate, La(ClO<sub>1</sub>)<sub>3</sub>, forms deliquescent needles, soluble

in alcohol (Cleve)

Lanthanum bromate, La(BrO<sub>3</sub>), 911,0 --- See p 256. When dried at 100°, this salt loses water and is converted into the dihydrate, La(BrO<sub>3</sub>)<sub>3</sub>.2H<sub>2</sub>O At 150° the anhydrous salt is obtained, and at high temperatures it readily decomposes

Lanthanum iodate, 2La(IO<sub>3</sub>)<sub>3</sub> 3H<sub>2</sub>O, obtained by double decomposition,<sup>8</sup> is sparingly soluble in water, one litre of a saturated solution at 25° containing  $1.87~{\rm grams}$  of anhydrous iodate  $^{\rm o}$ 

Lanthanum periodate, La10, 211,0, is obtained as a white, inicrocrystalline precipitate when aqueous solutions of lanthanum acetate and periodic acid are mixed (Cleve).

# LANTHANUM AND THE OXYGEN GROUP.

Lanthanum sesquioxide, or lanthana, La<sub>2</sub>O<sub>2</sub>, is obtained as a white powder by the ignition of the hydroxide, carbonate, nitrate oxalate, etc. The strongly ignited oxide has a density of 6.53 at 17° (Cleve), 5.94

Mangnac, Mem. Sci. phys. nat., 1855, 14, 201. (Euvies Completes (Geneva, 1902), vol. i. p. 362; Cleve, Bull. Soc. chim., 1874, [n.], 21, 196. Nechnesche, J. prakt. Chem., 1869, 107, 65; R. J. Meyer and Koss, Ber., 1902, 35, 2622.
 Mangnac, Euvies Completes (Geneva, 1902), vol. i. p. 640
 Alén, Oefreis Neuska Vet.-Akad. Förhandt., 1876, 33, Nos. 8, 9; Bull. Soc. chim., 1876, [ii.], 27, 365.
 Cleve, Ioc. ett.
 Frenchis and Smith, Annalen, 1878, 191, 331.
 Nilson, Ber., 1876, 9, 1056, 1142
 See p. 255, and Heimann, J. prakt. Chem., 1861, 82, 385, Frenchs, Ber., 1874, 7, 1798.

<sup>798.</sup>Holzmann, J. prakt. Chem., 1858, 75, 321, Heimann, thid, 1861, 82, 385.

Rimbach and Schubert, Zeitsch physikal Chem., 1909, 67, 183

(Hermann).1 It is slightly diamagnetic (see p. 257); the specific heat between 0° and 100° is 0.0749. The powder rapidly absorbs water, with the evolution of heat, and it also combines readily with carbon dioxide. The oxide crystallises in the form of hexagonal bipyramids of density 5:30 when dissolved in borax at a white heat and slowly cooled2, the crystals are not affected by water

Lanthana, even after intense ignition, is readily soluble in acids

Lanthanum hydroxide, La(OII), is produced by the direct union of lanthana and water, and may be obtained as a white, gelatmous precipitate by the addition of an excess of ammonia or alkali hydroxide to a solution of a lanthanum salt. The precipitate is sufficiently soluble in water to turn red htmus blue, it easily absorbs carbon dioxide from the air and sets free ammonia from ammonium salts. Lanthanum hydroxide is, in fact, the strongest base of all the rare earth hydroxides, being comparable with calcium hydroxide in its strength, and on this account the claim of Baskerville and Catlett to have prepared metallic lanthanates, in which lanthana acts as an acid, cannot be accepted until independent confirmation of their results is forthcoming.

Lanthanum peroxide -An oxide of lanthanum higher than La,Og cannot be prepared in the dry way, but indications of the existence of such an oxide have been obtained.

When a mixture of ammonia and hydrogen peroxide is added to a solution of a lanthanum salt, a gelatinous, hydrated perovide, La<sub>2</sub>O<sub>3</sub>xH<sub>2</sub>O, is obtained, to which the constitution (OH)<sub>2</sub>La(O OH) has been ascribed <sup>5</sup> Dilute sulphuric acid and carbonic acid set free hydrogen peroxide from the precipitate, and concentrated sulphuric acid decomposes it with the liberation of ozomsed

Lanthanum sulphide, LagS, the preparation and properties of which are given on p. 259, is a yellow solid of density 4 911 at 11" (Muthmann and

Stutzel)

Lanthanum persulphide, La<sub>2</sub>S<sub>p</sub>, is prepared by heating lanthanum sulphate to 580°-600' in hydrogen sulphide, and resembles the corresponding cerium compound in properties, decomposing into the sesquisulphide and sulphur at 650°.6

Lanthanum sulphite, La (SO), -A tetrahydrate of this salt has been

described by Cleve, and a tuhydrate by Grossmann (see p. 260)

Lanthanum sulphate,  $La_2(SO_1)_4$  — See p. 260 for general properties. Of all the rare earth sulphates, this is the most difficult to decompose completely by heat. Its specific heat (0 100 ) is 0 11827

One part of the anhydrons sulphate dissolves in 6 parts of ice-cold water, from the solution at 0° C, a hydrate, La<sub>2</sub>(SO<sub>4</sub>)<sub>2</sub>,1611<sub>2</sub>O, may be crystallised.5

Hermann, J. prakt. Chem., 1861, 82, 385
 Nordenskield, Porg. Annata., 1861, 114, 612, J. prakt. Chem., 1861, 85, 131.
 Baskerville and Catlett, J. Anna. Chem. Soc., 1994, 26, 75
 Marc, Ber., 1992, 2370, ef. Hermann, J. prakt. Chem., 1861, 82, 385, Zschiesche, ibid., 1869, 107, 65.
 Cleve, Bull. Soc. chim. 1885, [n.]. 43, 53, Melikoff and Pissarjewsky, Zeitsch. anorg Chem., 1899, 21, 70. See also Job. Compt. rend., 1993, 136, 45
 W. Biffz, Zeitsch. anorg. Chem., 1917, 127
 Nilson and Petterson, Ber., 1880, 13, 1459, Compt. rend., 1880, 91, 232.
 Brauner and Payliček, Trans. Chem. Soc., 1992, 81, 1243.

# ALUMINIUM AND ITS CONGENERS.

The hexagonal enneahydrate, La<sub>2</sub>(SO<sub>1</sub>), 9H<sub>2</sub>O, is the only hydrate stable in contact with the solution between 0° and 100°. It is readily prepared by warming a 15 per cent. ice-cold solution of lanthanum sulphate to 30°. The solubility diminishes with the temperature as follows:—2

Temp. ° C . . 0° 14° 18° 30° 50° 75° 100° Grams of 
$$\operatorname{La_2(SO_4)_3}$$
 3·0 2·6 2·1 1·9 1·6 1 0 0·7

The figures represent grams of anhydrous salt per 100 grams of water.

A hexahydrate, La<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>.6H<sub>2</sub>O, is said to crystallise from a solution in aqueous sulphuric acid (Frerichs and Smith).

The acid sulphate, La(HSO4), and the basic sulphate, La,O, SO, are known (p. 263). The following double salts have been described

```
\begin{array}{l} \operatorname{La_2(SO_4)_3.Cs_2SO_4.2H_2O}: ^6 \operatorname{3La_2(SO_4)_3.2Cs_2SO_1}^6 \\ \operatorname{La_2(SO_4)_3.Rb_2SO_4.2H_2O}: ^6 \operatorname{La_2(SO_4)_3.Rb_2SO_4}, ^6 \operatorname{3La_2(SO_4)_3.2Rb_2SO_4}^6 \\ \operatorname{La_2(SO_4)_3.K_2SO_4.2H_2O}: ^7 \operatorname{La_2(SO_4)_3.3K_2SO_4}, ^3 \operatorname{1} \\ \operatorname{La_2(SO_4)_3.5K_2SO_4}, ^7 \\ \operatorname{La_2(SO_4)_3.Na_2SO_1.2H_2O}.^{4.7} \\ \operatorname{La_2(SO_4)_3.Na_2SO_1.2H_2O}.^{4.5} \operatorname{with} \operatorname{2H_2O}, \operatorname{see} \operatorname{Barre}, ^7 \\ \operatorname{La_2(SO_4)_3.(NH_1)_2SO_4.7} \\ \operatorname{La_2(SO_4)_3.5(NH_1)_2SO_4}, ^7 \operatorname{La_2(SO_4)_3.5(NH_1)_2SO_4}, ^7 \operatorname{2La_2(SO_4)_3.5(NH_1)_2SO_4}, ^7 \end{array}
```

Lanthanum sulphate also combines with hydrazine sulphate 8

The diminution in solubility of lanthanum sulphate brought about by the addition of potassium, sodium, or ammonium sulphate will be seen from the following data, given by Barre, and denoting grams of anhydrous salts present in solution per 100 grams of water (solid phases not stated by Barre) -

| Temperature 16.5°.               |           | Temperature 18°.                  |                                                 | Temperature 18°.                                |            |
|----------------------------------|-----------|-----------------------------------|-------------------------------------------------|-------------------------------------------------|------------|
| K <sub>2</sub> SO <sub>4</sub> . | La2(SO4)3 | Na <sub>2</sub> SO <sub>4</sub> . | La <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> | (NH <sub>4</sub> ) <sub>2</sub> SO <sub>4</sub> | La2(SO4)3. |
| 0.00                             | 2:21      | 0 00                              | 2:13                                            | 0,00                                            | 2 13       |
| 0 25                             | 0.78      | 0.40                              | 1 00                                            | 4 01                                            | 0.39       |
| 0.50                             | 0.27      | 0.69                              | 0 35                                            | 8.73                                            | 0 28       |
| 0 85                             | 0 19      | 0 77                              | 0.30                                            | 18 24                                           | 0.25       |
| 1.03                             | 0 05      | 1 14                              | 0.13                                            | 27 89                                           | 0.48       |
| 1.16                             | 0.02      | 2 48                              | 0 04                                            | 36.11                                           | 0 28       |
| 2.50                             | 0 00      | 3 80                              | 0 02                                            | 47 49                                           | 0.14       |
|                                  |           | 5 55                              | 0.02                                            | 65 29                                           | 0.01       |

Lanthanum dithionate, La, (S2O6)3.24H2O, is readily soluble in water and orystallises in six-sided plates (Cleve).

3

<sup>1</sup> Kraus, Zeitsch. Kryst. Mun., 1901, 34, 307. See p. 261.

2 Muthmann and Rolig, Ber., 1898, 31, 1713; Barre, vide infra.

3 Hermann, J. prakt. Chem., 1861, 82, 385.

4 Cleve, Bull. Soc. chim., 1874, [in., 21, 196.

5 Marignac, Mem. Soc. phys. nat., 1855, 14, 201.

6 Baskerville and Moss, J. Amer. Chem. Soc., 1904, 26, 67.

7 Barre, Compt. rend., 1910, 151, 871.

8 Kolb, Zeitsch. anorg. Chem., 1908, 60, 123.

Lanthanum selenite,  $La_2(SeO_3)_3$ .—Hydrates of this salt with  $9H_2O$  and  $12H_2O$  have been described, and also the acid salts,  $La_2(SeO_3)_3$ . $2H_2SeO_3$ . $4H_2O$ and  $La_2(SeO_3)_3.3H_2Sc()_3.2H_2O^{-1}$ 

Lanthanum selenate, La2(SeO1)3.—Hydrates with 61120 and 10H2O have been described by Cleve, and also the following double salts: -

 $\begin{array}{l} {\rm La_2(SeO_1)_3.K_2SeO_1.9H_2O} \\ {\rm La_2(SeO_1)_3.Na_2SeO_1.4H_2O} \end{array}$ 

 $\text{Lsi}_{2}(\text{SeO}_{4})_{3} (\text{NH}_{1})_{2}\text{SeO}_{1}.9\text{H}_{2}\text{O}$ 

Lanthanum chromate, Lag(C1O4), 8H2O, may be obtained by double decomposition as a sparingly soluble, yellow, crystalline precipitate (Clove).

Lanthanum molybdate, Lag(MoO<sub>1</sub>), has been prepared (p. 265). Lanthanum tungstate, La<sub>1</sub>(WO<sub>1</sub>)<sub>3</sub>, the metatungstate, double and complex tungstates have been described (p. 265)

Lanthanum silicotungstate, La<sub>1</sub>(W<sub>12</sub>SiO<sub>10</sub>),—See p 266.

### LANTHANUM AND THE NITROGEN GROUP.

Lanthanum nitride, LaN, has been described (p. 267)

Lanthanum azide (hydrazoute, trinitride).—The basic salt 21a(N3)2(OH).

3H2O is known and is very explosive.2

Lanthanum nitrate, La(NO<sub>3</sub>), 6H<sub>2</sub>O, forms deliquescent, triclinic crystals <sup>3</sup> which lose 5H<sub>2</sub>O over sulphuric acid. <sup>1</sup> At 25" a saturated aqueous solution of the hexahydrate contains 151 1 parts of anhydrous salt per 100 of water. Lanthanum nitrate is isodimorphous with bismuth intrate (see p. 234).

Lanthanum ammonium nitrate, La(NO3), 2(NH1)NO3 1H2O, has already

been described.6

Lanthanum cassum netrate, La(NO<sub>2</sub>), 2CsNO<sub>1</sub> 2H<sub>2</sub>O<sub>2</sub>, forms monochnic crystals (a . b : c = 1 3052 | 1 | 0 9663 | B = 103° 26′). T

Lanthanum rubidium nitrate, La(NO<sub>3</sub>), 2RbNO<sub>3</sub> 4H<sub>2</sub>O, has been described

Lanthanum potassium nitrate, La(NO<sub>3</sub>), 2KNO<sub>2</sub>H<sub>2</sub>O, forms orthorhombic crystals (hemimorphic, a·b.c=05306 1 0·5696) s of density 2·54 at 0° C.

Lanthanum sodium nitrate, La(NO<sub>3</sub>), 2NaNO<sub>3</sub>H<sub>2</sub>O, forms monoclinic crystals (a·b.c=19970 1.07678; β=90° 32°) of density 2·63 at 0° C s

Lanthanum thallium nitrate, La(NO3)3 2TINO1.4H,O, has been described (p. 268).

Acid lanthanum rubidium nitrate, La(NO<sub>3</sub>), RbNO<sub>3</sub>, HNO<sub>4</sub> 6H<sub>2</sub>O, has been prepared by Jantsch and Wigdorow. When heated to 120° it loses nitric acid and water, leaving the salt La(NO<sub>3</sub>), RbNO<sub>4</sub> H<sub>2</sub>O.

Cleve, loc. cit., Nilson, Nova Lite Soc Uprata, 1875, [in ], 9, No. 7, Ball, Soc chim.,
 Cartins and Datapsky, J. pratt. Chem., 1900, [ii ], 61, 108
 See p. 268; and Marignac, Ann. Chim. Phys., 1873, [iv.], 30, 56.
 Mosander, Phil. May, 1843, [iii], 23, 241, Cleve, Ball. Soc. chim., 1874, [ii.], 21, 196; Hermann, J. prakt. Chem., 1861, 82, 385; Zschnesche, bid., 1°09, 107, 65
 James and Whittemore, J. Amer. Chem. Soc., 1912, 34, 1168.
 See p. 268; Marignac, loc. cit., 1906t, Bull. Soc. franç. Min., 1888, 11, 143, 215;
 Kraus, Zeitsch. Kryst. Min., 1901, 34, 307
 Wyrouboff, Bull. Soc. franç. Min., 1907, 30, 299, Jantsch and Wigdorow, Zeitsch. anorg. Chem., 1911, 69, 221.
 Fock, Zeitsch. Kryst. Min., 1894, 22, 37, Wyrouboff, Bull. Soc. franç. Min., 1907, 30, 299; Jantsch and Wigdorow. loc. cit.
 Wyrouboff, loc. cit., Jantsch and Wigdorow, loc. ck.

For the double nitrates of lanthanum and bivalent metals, 3M11 (NO<sub>3</sub>)<sub>2</sub>. 2La(NO<sub>3</sub>), 24H<sub>2</sub>O, see p 269.1

Lanthanum phosphite, I.a. (HPO,), has been described by Frerichs

and Smith.

Lanthanum orthophosphate, LaPO<sub>1</sub>, is obtained as a white, amorphous precipitate by adding phosphoric acid or sodium orthophosphate to a solution of a lanthanum salt (Cleve)—It may be obtained in the crystalline form by heating lanthana with molten potassium metaphosphate.<sup>2</sup> The double salt 2LaPO<sub>4</sub> K<sub>1</sub>PO<sub>4</sub> may be prepared by saturating molten potassium pyro- or ortho-phosphate with lanthana.

Lanthanum hydrogen pyrophosphate, LaHP<sub>2</sub>O<sub>7</sub>.3H<sub>2</sub>O —From a solution of a lanthanum salt sodium pyrophosphate throws down a white precipitate which then redissolves On allowing the solution to stand, the

acid pyrophosphate crystallises out in needles (Cleve)

The double salt LaNaP 07 may be obtained in crystals by fusing together

sodium metaphosphate and lanthana

Lanthanum metaphosphate, La(PO<sub>3</sub>), and the compound La<sub>2</sub>O<sub>3</sub>,5P<sub>2</sub>O<sub>5</sub>

have been prepared.4

Lanthanum arsenite, Lag(IIAsO3), and lanthanum arsenate, Lag(HAsO1), have also been described (Frerichs and Smith)

### LANTHANUM AND THE CARBON GROUP.

Lanthanum carbide, LaC, has been described (p. 270) Lanthanum carbonate, La<sub>2</sub>( $(O_n)_1$ 811,0, loses 711,0 at 100°. The octahydrate occurs in nature as the orthorhombic mineral lanthanite (a. b. c=-0.9528 + 1 + 0.9023) The following double salts are known --5

$$\begin{array}{lll} \operatorname{Lat_2(CO_3)_3} & \operatorname{K_2(CO_3, 12H_2O} & 2\operatorname{Lat_2(CO_3)_3} & \operatorname{3Na_2CO_3, 20H_2O} \\ \operatorname{Lat_2(CO_3)_3} & \operatorname{NH_1)_2(O_3, 4H_2O} & \end{array}$$

Lanthanum thiocyanate, La(CNS), 7H<sub>2</sub>O, and the double salt, La(CNS), 3Hg(CN)<sub>2</sub>.12H<sub>2</sub>O, are known (Cleve, see p. 272)

Lanthanum platinocyanide, 2La(CN), 3Pt(CN2, 18H2O, forms yellow

prisms having a blue reflex (Cleve, see p. 272).

Lanthanum acetate, 2La(C<sub>2</sub>H<sub>1</sub>O<sub>2</sub>), 3H<sub>2</sub>O, crystallises from its aqueous

solution in thin prisms.6

When a cold aqueous solution of lanthanum acetate is made ammoniacal, a colloidal basic acetate is precipitated; if a little solid iodine be added to the precipitate a blue adsorption compound is produced, similar in appearance to the familiar "starch-iodide" precipitate.7 The blue substance may conveniently be prepared by adding a solution of iodine in potassium iodide

Also Holzmann, J. prakt. Chem., 1858, 75, 321, Carius, whil, 1858, 75, 352; Grant and James, J. Amer. Chem. Soc., 1915, 37, 2652 (Fe)
 Ouvrard, Compt. rend., 1888, 107, 37, Grandeau, Ann. Chim. Phys., 1886, [vi.], 8, 166. Ouvrard, Compt. rena., 2-1.
193.
Wallroth, Bull. Soc chim, 1883, [ii.], 39, 316
Frenichs and Smith, loc cit., Johnson, Ber., 1889, 22, 976
R. J. Meyer, Zev'sch. anorg. Chem., 1904, 41, 97, see p. 271.
Czudnowicz, J. publ. ('hem., 1880, 30, 31; Cleve, Bull. Soc. chim., 1871, [ii.], 21, 196; Behrens, Arch. Néerland., 1991, [ii.], 6, 67.
Damour, Compt. rend., 1856, 43, 976.

to a cold solution of lanthanum acctate (or nitrate acidified with acetic acid), adding ammonia cautiously until very little iodine is left unchanged, and then warming the mixture very gently <sup>1</sup>

Lanthanum ethylsulphate, La(C2H3-SO1), 9H2O. See p 278.

Lanthanum acetylacetonate, La(CII, CO CII (O CII,), has also been

described (p. 279)

Lanthanum oxalate, La<sub>2</sub>(C<sub>2</sub>O<sub>4</sub>), 9(10 or 11)H<sub>2</sub>O<sub>5</sub><sup>2</sup> may be obtained by double decomposition as a white, crystalline precipitate. According to the temperature at which precipitation occurs and the temperature of the wash water, the precipitate appears to contain 9, 10, or 11H<sub>2</sub>O. Most experimenters give 10H<sub>2</sub>O as being present in the hydrate in equilibrium with its aqueous solution at 25°.

Lanthanum oxalate is appreciably soluble in a solution of lanthanum intrate, but no crystalline oxalonitrate can be isolated at 25°. The only hydrates of lanthanum oxalate that are stable at 25° in contact with aqueous solutions of lanthanum intrate of the appropriate concentrations are those containing 3H<sub>2</sub>O, 5H<sub>2</sub>O, and 8H<sub>2</sub>O<sup>-3</sup>. A hydrate containing 7H<sub>2</sub>O may be crystallised from dilute sulphunic acid <sup>4</sup>

A solution of lanthanum oxalate in hot, concentrated hydrochloric acid deposits crystals of lanthanum oxalothloride, 2La(C<sub>2</sub>O<sub>4</sub>)Cl 511<sub>2</sub>O, when cooled

The salt is decomposed by water '

For further details concerning lanthanum oxalate, see p. 273

Lanthanum salts of organic acids—The following salts have been described—

Lanthanum formate, b succenate, b, 7, 8, 9 tartrate, b, 7, 10, 12 citrate, 7, 12 schacate, 17 glycollate, 13 mulomate, 11, 12 demethylphosphate, 19 henroate, 7 phenylacetate, 13 hyppurate, 7 phenoxyacetate, 16 henroa calphonate, 12 m actio, methodoo, and m-homobenrenessulphonate, 12 p-dichlorobenrenessulphonate, 14 for 3, 1-thlorometrobenrenessulphonate, 14 for 3, 1-thlorometrobenrenessulphonate, 18 8-hydroxynaphthalene-1-sulphonate, 11 a-naphthalenesulphonate, 12 for 7-nitronaphthalenesulphonate 12 and pyromucate 20

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1 W. Biltz, Bee, 1904, 37, 719.
2 9H<sub>2</sub>O, Crudinowice, J. peak. Chem., 1860, 80, 31. Cleve, Bull. Soc. chem., 1874, [1], 21, 196. 10H<sub>2</sub>O, Power and Shedden, J. Soc. Chem. Ind., 1900, 19, 646, Hauser and Witth, Zeitsch. anal. Chem., 1908, 47, 389, Witth, Zeitsch. anang. Chem., 1912, 76, 174. 11H<sub>2</sub>O, Wyrouboll. Bull. Soc. trans. Um., 1901, 24, 105; Brauner and Pavlièsk, Trans. Chem. Soc., 1902, 81, 1213
3 Whitteniore and James. J. Amer. Chem. Soc., 1912, 34, 1168.
1 See Witth, Zeitsch. anang. Chem., 1908, 58, 213
5 Job. Compt. rend., 1898, 126, 219
6 Cleve, Bull. Soc. chem., 1874, [1], 21, 196.
7 Caudnowicz, J. prokt. Chem., 1800, 80, 31.
8 Behrens, Arch. Nerdand, 1901, [1], [6, 67]
9 R. J. Meyer, Zeitsch. anang. Chem., 1902, 33, 113
10 Rimbach and Schubert, Zeitsch. physikat. Chem., 1909, 67, 183.
11 Erdmann and Witth, Janalea, 1908, 367, 190
12 Holmberg, Zeitsch. anang. Chem., 1907, 53, 83.
13 Jantsch and Giunkraut, ibid., 1913, 79, 305
14 Armstrong and Rodd, Proc. Roy. Soc., 1912, A, 87, 204.
15 Rodd, ibid., 1913, A, 89, 292
16 Pratt and James. J. Amer. Chem. Soc., 1911, 33, 1330.
17 Whittemore and James, ibid., 1913, 35, 127.
18 Katz and James, ibid., 1913, 35, 57;
19 Morgan and James, ibid., 1913, 37, 2652.
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#### LANTHANUM AND BORON.

Lanthanum metaborate, La(BO<sub>2</sub>)<sub>3</sub>, has been described (p 282).

### PRASEODYMIUM.

Symbol, P1. Atomic weight, 140.921 (0=16).

Praseodymium is one of the most abundant of the rare earth elements, but it is decidedly less plentiful than lanthanum or needymium. The relative proportions of prascodymium and neodymium in different minerals do not vary very much, the ratio praseodymium to neodymium being approximately 1 to 2.2

The occurrence, history, preparation, atomic weight, and homogeneity of praseodymium have been already discussed in Chapters X and XI.

Praseodymium 3 is a yellow metal of density 6:475 It melts at 940° and is strongly paramagnetic, the magnetic susceptibility at 18' being about  $25 \times 10^{-6}$  c.g s. units per gram. Very little else is known of the properties of the metal.

# COMPOUNDS OF PRASEODYMIUM.

The salts of prascodymium are derived from the basic oxide Pr<sub>2</sub>O<sub>3</sub>. They are leek-green in colour, and their reflection spectra or the absorption spectra of their solutions exhibit characteristic absorption bands. The conductivities of prascodymium salts in aqueous solution are in harmony with the view that the salts are derived from a fairly strong triacid base. The following results, for example, refer to the chloride at  $25^{\circ}$  (v=dilution in litres per gram equivalent) -5

Praseodymium is of interest in that it forms a dioxide, PiO, According to Brauner, the dioxide is a weak base, and gives rise to a basic sulphate,  $2Pr_2O_1.SO_3 + 2\frac{1}{2}H_2O$ , and a basic acetate,  $(CH_3.CO)_3.PrO.Pr(OH)_3$  or  $\begin{array}{l} (CH_3\cdot CO)_2\cdot \Pr_2O_2(OH)(CO.CH_3) + II_2O.^6 & \text{The same authority also mentions an acetate of the constitution } (CH_3\cdot CO)_2\cdot \Pr_1O\cdot O.O\cdot \Pr_1(OII)(CII_1\cdot CO) + H_2O.^7. \end{array}$ 

Numerous compounds of praseodymnum have been described by von Schéele.8

Thermochemistry of Praseodymium.—The following results have been obtained .- 9

<sup>1</sup> Bayter and O. J. Stewart. J Amer. Chem. Soc., 1915, 37, 516.
2 Muthmann and Stutzel, Ber., 1899, 32, 2853.
3 Muthmann and Weiss, Annalen, 1904, 331, 1. See also pp. 229-230.
4 Owen, Ann Physik, 1912, [iv.], 37, 657.
5 Aufrecht, Inaugural Dissertation (Berlin, 1904); see also Jones and Reese, Amer. Chem. J., 1898, 20, 606.
6 Brauner, Proc. Chem. Soc., 1898, 14, 70.
7 Brauner, Proc. Chem. Soc., 1901, 17, 66.
7 Von Schéele, Zeitsch. anorg. Chem., 1898, 17, 310; 18, 352
8 Muthmann and Weiss, Annalen, 1904, 331, 1 (2Pr+30); Matignon, Ann. Chim.: Phys., 1906, [viii], 8, 386; Compt. rend., 1906, 142, 276.

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\begin{array}{lll} 2[\Pr] + 3(O) & = \Pr_2O_3 \\ [\Pr_2O_3] + 6HClaq. & = 2\Pr_2Cl_3aq. + 3H_2O \\ [\Pr_2O_3] + 6(HCl) & = 2[\Pr_2Cl_3] + 3[H_2O] \\ [\Pr_2O_3] + 3[H_2SO_1] = [\Pr_2(SO_1)_3] + 3[H_2O] \\ [\Pr_2Cl_3] + \Delta q. & = \Pr_2Cl_3aq. \\ [\Pr_2Cl_3] + 2O[A_2] + \Delta q. & = \Pr_2Cl_3aq. \\ [\Pr_2Cl_3] + 2O[A_2] + \Delta q. & = \Pr_2Cl_3aq. \end{array}
2|Pr| + 3(0)
   +412.4 Cals.
  +1062
   +147.8
  +1257
  +335
   28 9
   [PrCl_3^*7H_2O] + Aq. = PrCl_3Aq.
  53
   \Pr_2(\mathrm{SO}_4)_3aq
     \Pr_2(SO_4)_3] + Aq.
   36.0
```

Praseodymium hydride, PrII, has been described (p. 251).

### PRASEODYMIUM AND THE FLLORINE GROUP.

Praseodymium fluoride, PrF, prepared from aqueous praseodymium sulphate and hydrofluoric acid, is a yellow precipitate. When dried, it forms

glittering yellow crystals which appear green by reflection <sup>1</sup>

Praseodymium chloride, <sup>2</sup> PrG, —The methods of preparation and properties of the anhydrous salt are given on p 252. It is a green, hygroscopic solid which fuses to a green liquid. The density is 4.020 at 25° C. (Baxter and Stewart).

From aqueous solution the heptahydrate, PiCl, 711,0, separates in large, green crystals of density 2.25 at 15° (Scheele) The crystals melt at 111° C. (Baxter and Stewart). When dued over sulphure acid, the heptahydrate rapidly passes into the hexahydrate, PiCl, 6H2O, which is slowly dehydrated to the trihydrate, PrCl, 3H,0. The monohydrate is also known (p. 254). At 13 8°, 100 parts of water dissolve 103 9 parts of anhydrous chloride or 334.2 of the heptahydrate, the solution being in equilibrium with the heptahydrate (Matignon). The concentrated aqueous solution dissolves considerable quantities of rare earth ovalates.

Prascodymium aurichlorule, Pr(II, AuCl3.101120, forms yellow crystals of specific gravity 2 60 and is readily soluble in water. The chloroplatinate, PrCl. PtCl. 12H.O, is also a yellow crystalline salt, of density 2 41 at 16°.

Praseodymium oxychloride, PrOCl, has been prepared (p. 255). Praseodymium bromide, PrBr,—The anhydrous salt has been prepared, and closely resembles the chloride (p. 255). The hoxahydrate, PrBr, 6H<sub>2</sub>O, crystallises from aqueous solution (von Schéele).

Praseodymium iodide, Pri, has been prepared by Matignon (see p. 255). Praseodymium bromate, Pr(Br(),), 9H2O, forms green hexagonal prisms (p. 256). It is converted into the dihydrate, Pr(BrO<sub>3</sub>)<sub>3</sub>.2H<sub>2</sub>O, at 100° and the anhydrous salt at 130°, while it decomposes at 150°.

# PRASEODYMIUM AND THE OXYGEN GROUP

Praseodymium sesquioxide, or praseodymia, Pr<sub>2</sub>O<sub>g</sub> is obtained by reducing the dioxide at a red heat in a stream of hydrogen. It is a yellowishgreen powder of density 6 883 (or 7 074) at 15°. When heated in the air it absorbs oxygen and becomes brown in colour.

<sup>&</sup>lt;sup>1</sup> Popovici, Ber , 1908, 41, 634. Matigpon, Compt rend, 1902, 134, 427; Ann Chim. Phys. 1908, [vii.], 8, 384;
 Bourion, obid., 1910, [viii.], 21, 49; of R. J. Meyer and Koss, Err., 1902, 35, 2622;
 Barter and Stewart, J. Amer. Chem. Soc., 1915, 37, 516.
 Von Schéele, Zeitsch. anorg. Chem., 1898, 17, 310.
 Brauner, Proc. Chem. Soc., 1901, 17, 66.

Praseodymium hydroxide, Pr(OH), may be obtained by double decomposition as a bright green, gelatinous precipitate. In the presence of hydrogen perovide, ammonia produces with prascodymium salts a precipitate of hydrated mascodymum peroxule, Pr.O5 xH2O or (OH), Pr O OH. The peroxide closely resembles the lanthanum compound in its properties 1

Praseodymium dioxide, PrO, may be prepared by heating the nitrate to 410° or by fusing the intrate with intre at 400° until decomposition is complete 2. It is a black solid of density 5 978 at 20°. According to Brauner, the pure oxide is not of the hydrogen peroxide type, but contains quadrivalent praseodymium. It sets free chlorine from hydrochloric acid, iodine from hydrodic acid, and ozonised oxygen with oxyacids 3 it oxidises cerous salts from coric, manganous salts to permanganic acid, and gives a violet coloration with a solution of strychime in sulphuric acid. It is reduced by hydrogen peroxide in acid solution. It oxidises ferrous and stannous salts, but a part of the available oxygen is always lost (von Schéele).

When praseodymum hydroxide, carbonate, intrate, oxalate, etc., are heated, dark brown oxide residues are obtained which are intermediate in composition between  $Pr_2O_3$  and  $PrO_9$  and which contain less oxygen in proportion as the temperature at which they are produced increases antermediate oxides of the formula  $\text{Pr}_1\text{O}_{7,5}$   $\text{Pr}_2\text{O}_{9,6}$  and  $\text{Pr}_4\text{O}_{1,7}$  have been described, but their individuality is doubtful. Brauner considers the oxide  $Pr_5O_9$  to be a complex oxide formed by the combination of  $Pr_2O_1$  with  $PrO_2$   $(Pr_5O_9=Pr_2O_9,3PrO_9)$  . The formation of praseodymium dioxide in a mixture of cerumi earths

is favourably influenced by the presence of cerra, which probably acts as an oxygen-carrier, but is adversely affected by the presence of lanthana or ncodyma.8

Praseodymium sulphide, Pr.S., cannot be prepared by the usual methods (see p. 259) if entirely free from cornum, the sulphate, when heated in hydrogen sulphide, giving only prascodymium oxysulphide, Pr<sub>2</sub>O<sub>2</sub>S. If, however, the prascodymium sulphate contains a little cerous sulphate, the trisulphide, Pr<sub>2</sub>S<sub>3</sub> and persulphide, Pr<sub>2</sub>S<sub>3</sub>, may be obtained as in the cases of cerium and lanthanum. The trisulphide is chocolate-brown in colour and has a density of 5 012 at 11° (Muthmann and Stutzel). The persulphide begins to lose sulphur at about 650° 9

Praseodymium sulphate, Pr<sub>2</sub>(SO<sub>1</sub>)<sub>4</sub>.—For preparation and general properties, see p. 260 At 0°, 23 6 parts, and at 20, 17·7 parts of the anhydrous sulphate can be dissolved in 100 of water

The monoclime  $^{10}$  octahydrate,  $\Pr_2(\mathrm{SO}_1)$ ,  $8\mathrm{H}_2\mathrm{O}_1$  is the only hydrate stable in

Mehkoff and Khmenko, J. Russ. Phys. Chem. Soc., 1901, 33, 503
 Brauner, Proc. Chem. Soc., 1898, 14, 70., 1901, 17, 66
 See, however, Mehkoff and Khmenko, J. Russ. Phys. Chem. Soc., 1901, 33, 739.
 Von Scheele, Inc. ett.
 Arer von Welsbech, Monatsh., 1885, 6, 477.
 Brauner, Proc. Chem. Soc., 1898, 14, 70., 1901, 17, 66.
 Schottlander, Ber., 1892, 25, 569, R. J. Meyer, Zeitsch. among. Chem., 1904, 42, 47

<sup>41, 97.</sup>Brauner, Monatsh , 1882, 3, 1; Maic, Ber , 1902, 35, 2370 , R. J. Meyer and Koss,

Ber., 1902, 35, 3740.

W. Biltz, Zeitsch. anorg. (ham., 1911, 71, 427.

Kraus, Zeitsch. Kryst. Min., 1901, 34, 307; Dufet, Cham. Zento., 1902, 1, 452. See p. 261.

contact with the solution from 0°-75°; 1 its solubility, in grains of anhydrous salt per 100 grams of water, is as follows :-

Temp 'C . . . 0' 18' 35' 55' 75° Grains of 
$$Pr_2(SO_4)_3$$
 . 19 80 14 05 10 40 7 02 1 20

The monoclinic 2 pentahydrate, Pr. (SO<sub>4</sub>), 511,0, separates from the aqueous solution at 90 - 100 . Its solubility, in grains of anhydrous sulphate per 100 grams of water, is as follows (Muthmann and Rolig) - -

The transition point (from octa- to penta hydrate) is about 75 Two labile hydrates, Pr<sub>2</sub>(8O<sub>4</sub>), 154H<sub>2</sub>O and Pr<sub>2</sub>(8O<sub>4</sub>) 12H<sub>2</sub>O, have been isolated at low temperatures by von Schoole and by Mithmann and Rolig respectively; Braimer mentions a hexahydrate \*

For acid and basic sulphates, Pr(HSO<sub>4</sub>), and Pr<sub>2</sub>O<sub>5</sub>SO<sub>5</sub> respectively, see p 263
The following double sulphates have been described

```
\begin{array}{l} \Pr_1(80_1), (S, 80_1, 2H, 0.^5 \text{ and } \Pr_2(80_1), (S, 80_1, 4H, 0.^5 \\ \Pr_2(80_1), (K, 80_1, 4H, 0.^5, P_1, (80_1), 3K, 80_1, H, 0.^4 \\ \text{density } 3/29 \end{array}
Pr<sub>2</sub>(SO<sub>1</sub>)<sub>r</sub>(NH<sub>1</sub>)<sub>2</sub>SO<sub>1</sub> 8H<sub>2</sub>O<sub>2</sub> density 2.53
```

Praseodymium dithionate, Pr<sub>2</sub>(S<sub>2</sub>O<sub>6</sub>), 1211,O, is extremely soluble in water (von Scheele)

Praseodymium selenite - The basic salt precipitated by sodium sclenite from prascodymium sulphate solution is converted into the acid selenite, Pr. (SeO<sub>3</sub>), H<sub>2</sub>SeO<sub>3</sub> 3H<sub>2</sub>O<sub>5</sub> by heating with aqueous selenions acid (von Schéele)

Praseodymium selenate,  $Pr.(SeO_1)_3$ —The octahydrate,  $Pr.(SeO_1)_3$ 811,0, crystallises from cold aqueous solutions, and the pentahydrate, Pr. (SeO4), 511, O, from hot solutions The double salt prascodymium potassium sitenate,  $\Pr_{u}(SeO_1)$ ,  $3K_uSeO_1$  III,O, is somewhat more soluble than the double sulphate (von Scheele)

Praseodymium molybdate, Pr.(MoO<sub>1</sub>) - See p. 265

Praseodymium tungstate, Pr. (WO<sub>1</sub>), and complex tungstates have been already mentioned (p 265)

Praseodymium silicotungstate, Pr<sub>i</sub>(W<sub>10</sub>SiO<sub>m</sub>) See p. 266

### PRASEODYMILM AND THE NITROGEN GROUP

Praseodymium nitride, P(N) has been described (p. 267). Praseodymium nitrate,  $P(NO_D)$  6H<sub>2</sub>O, crystallises in large, green needles The following double salts are known Pr(NO<sub>3</sub>), 2(NH<sub>1</sub>)NO<sub>3</sub> 4H<sub>2</sub>O; Pr(NO<sub>3</sub>)<sub>3</sub> 2RbNO<sub>3</sub> 4H<sub>2</sub>O<sub>4</sub>, Pr(NO<sub>3</sub>)<sub>4</sub> 2NaNO<sub>4</sub> H<sub>2</sub>O (von Scheele) and the double salts with the intrates of bivalent metals, 3Mn(NO<sub>3</sub>), 2Pr(NO<sub>3</sub>), 21H<sub>2</sub>O (see p 268)

<sup>&</sup>lt;sup>1</sup> Muthmann and Rolig, Iter 1898, 31, 1718.

<sup>2</sup> Krans, los ett See p 261 4 Brauner, Proc Chem Sec 1898, 14 70 4 Von Schiele, los ett

Baskerville and Holland, J. Amer. Chelp Soc., 1901, 26. 71.

# PRASEODYMIUM AND THE CARBON GROUP.

Praseodymium carbide, PrC<sub>2</sub>, has been described (p. 270). Praseodymium carbonate, Pr<sub>2</sub>(CO<sub>3</sub>)<sub>4</sub>,8H<sub>2</sub>O, has been prepared by von Scheele. The following double salts are known :----1

> $\begin{array}{l} \Pr_2(\text{CO}_3)_3 \; \text{K}_2\text{CO}_3.12 \, \text{H}_2\text{O} \\ \Pr_2(\text{CO}_3)_3 \; (\text{NH}_1)_2\text{CO}_3 \; \text{1H}_2\text{O} \end{array}$ 2Pr<sub>2</sub>(CO<sub>3</sub>)<sub>3</sub> 3Na<sub>2</sub>CO<sub>3</sub>·22H<sub>2</sub>O

Praseodymium ethylsulphate, Pr(C2H3SO1), 9H2O, has been described (p. 278).

Praseodymium acetylacetonate, Pi(CH<sub>3</sub> CO.CH CO CH<sub>3</sub>)<sub>3</sub>, has also been described (p 279).

Praseodymium oxalate, Pr<sub>2</sub>(C<sub>2</sub>O<sub>1</sub>), 10H<sub>2</sub>O, may be obtained by double decomposition as a green, crystalline precipitate. For its solubility in water and acids, see p. 273

Praseodymium salts of organic acids. The following salts have been described -

Praseodymum acctate, 2 propionate, 2 malonate, 10 citrate, 5 succinate, 3,4 sebacate, 7  $gly collate, ^{12} car only late, ^{7} demothyl phosphate, ^{9} phonory acctate, ^{6} benzene sulphonate, ^{11} m-chlor obenzene sulphonate, ^{11} 6:3 1-chlor on tro$ benzenesulphonate, 11 1 4 · 2-bromonitrobenzenesulphonate, 8 p-dichlorobenzenesulphonate, 11 p-dibromobenzenesulphonate, 13 a-naphthalenesulphonate, 11 1:5-, 1:6-, and 1 7-nitronaphthalenesulphonate, 11 8-hydroxynaphthalenes 1-sulphonate 10

Basic mascodymium acetate gives rise to a violet-blue absorption compound with rodine in a similar manner to the lanthanum compound (p. 410) 15

# NEODYMIUM.

Symbol, Nd. Atomic weight, 144.3 (0=16)

Neodymum is one of the most abundant of the rare earth elements, being about twice as plentiful as prascodymium (p. 112). Its occurrence, history, preparation, atomic weight, and homogeneity have been already discussed in Chapters X and XI

• Neodymium is a pale yellow metal of density 6.956. It melts at 840°. Neodymium is decidedly paramagnetic, the magnetic susceptibility at 18° being about  $36 \times 10^{-6}$  c g s units per gram <sup>17</sup>

```
1 R. J. Meyer, Zeitsch. anory Chem., 1904, 41, 97. see p. 271.
2 Von Schiede, Zeitsch. anory Chem., 1898, 17. 310. 18, 352.
3 Behrens, Arch. Nerland., 1901, [u] 6, 67.
4 R. J. Meyer, Zeitsch. anorg. Chem., 1902, 33, 113.
5 Baskenville and Turrentine, J. Amer. Chem. Soc., 1904, 26, 46.
6 Pratt and James, ibid., 1911, 33, 1330.
7 Whittemore and James, ibid., 1913, 35, 872.
8 Katz and James, ibid., 1914, 36, 10.
10 Erdmann and With, Annalen, 1908, 361, 190.
11 Holmberg, Zeitsch. anorg. Chem., 1907, 53, 83.
12 Jantsch and Grunkmut, ibid., 1918, 79, 305.
13 Armstrong and Rodd, Proc. Roy. Soc., 1912, A, 87, 204.
14 Rodd, ibid., 1913, A, 89, 292.
15 Oloff, Chem. Zeit., 1907, 31, 45.
16 Muthmann, Hofer, and Weiss, Annalen, 1902, 320, 231, Muthmann and Weiss, ibid., 1904, 331, 1; Muthmann and Scheidmandel, ibid., 1907, 355, 116. See pp. 229-230.
17 Owen, Ann. Physik, 1912, [v.], 37, 657.
```

# COMPOUNDS OF NEODYMUM

The salts of noodymum are rose red or reddish-violet in colour, and the absorption spectra of their aqueous solutions show characteristic absorption bands. The following equivalent conductivity determinations ( $\lambda$ ), effected at 25 , show that the salts are derived from a fairly strong trucid base ( r =dilution in litres per grain equivalent)

```
936.96
NdCL
                  29.28
                            58.56 - 117.12
   231 21 468 18
   1211
   128.7
  133 4
             \lambda = 103.0
                           1109
                                    118.85
                                    133.88
   267 76 535 52
  1071 01
Nd (801)1
                  33 17
                            66.91
  95 95
   80.21
                 10.61
                            18 15
                                      17 12(4) 62 23
                            67.62
                                    135.21
   270.48
  -540.96
   1081.92
\mathrm{Nd}(\mathrm{C}_2\mathrm{H},\mathrm{O}_2)/\tau = -33.81
   86.89
   67.66
   76.81
                            50.28
                                     58 55
             λ. 12 Β
```

Thermochemistry of Neodymium. The following results have been recorded - -2

| 2[Nd]+3(O)                                                         | $[Nd_iO_i]$ .                         | + 135 1 Cals | , |
|--------------------------------------------------------------------|---------------------------------------|--------------|---|
| 2, Nd] + 3[S]                                                      | [Nd 8.]                               | 4 285 9 .,   |   |
| [Nd]+3(Cl)                                                         | - [NdCl]                              | + 2195       |   |
| [Nd] i 3[1]                                                        | [NdL]                                 | +1577 ,,     |   |
| $2[Nd] + 3[S] + 6(O_0)$                                            | $\{Xd_i\hat{S}O_D\}$                  | + 925 1 ,,   |   |
| $2[Nd] + 3[S] + 6(O_0) + 8[H.C]$                                   |                                       | 1 916 8      |   |
| $[NdCl_i] + Aq$                                                    | `\dClaq`                              | + 354        |   |
| NdC1 6H <sub>2</sub> O]+ Aq                                        | NdCLaq                                | + 76         |   |
| $[NdL] + \sqrt{q}$                                                 | NdLaq                                 | 1 189        |   |
| $[Nd_{s}(SO_{1})_{s}] + \Lambda q$                                 | Nd (SO <sub>c</sub> ) aq              | 1 36 5 ,.    |   |
| $[Nd.(SO_i), 5H_iO] + \lambda q$ .                                 | Nd (80) aq                            | 4 83 ,,      |   |
| $N_{\bullet}$ (SO <sub>3</sub> ) 8H <sub>2</sub> O + $\Lambda_{0}$ | Nd,(80,)q                             | + 67 ,,      |   |
| NdCl.] + alcohol                                                   | = NdCl, m alc soln                    | 1 21 5 ,     |   |
| $[Nd_2O_3] + 6HCLq$                                                | - 2NdCl aq + 3H,O                     | + 105.5 ,,   |   |
| Nd.O.] + 6111aq                                                    | $2NdLaq + 3H_3O$                      | 11061 ,,     |   |
| Nd.o.1 + 3fH.So.1                                                  | $Xd_{2}(SO_{1})_{3}] + 3[H_{2}O_{1}]$ | +125:1       |   |
| \d]0] + 3[H,\$0]<br>\Xd_0] + 3H,\$0pq                              | $Nd_{2}(SO_{1})$ aq $\pm 3H_{2}O_{1}$ | + 106 1 ,,   |   |
|                                                                    |                                       |              |   |

### NEODYMEM AND HYDROGEN

Neodymium hydride, NdH, has been already described (p. 251).

# NEODYMUM AND THE FITORINE GROUP

Neodymium fluoride, NdF, from aqueous neodymium sulphate and hydrofluoric acid, is deposited on warning as a pale like coloured crystallino powder, insoluble in water 4

Neodymium chloride, AdCl —For the preputation and properties of the anhydrous salt, see p. 252. It is a hygroscopic, rose-coloured, crystalline mass of density 4434 at 25. (Baxter and Chapin).

<sup>&</sup>lt;sup>1</sup> Autrocht, Inaugural Description (Berlin 1904 <sup>2</sup> Matignon, Compt. read., 1906, 142, 276. Ann. Chem. Phus., 1906, [viii.] 8, 213., 1907, [viii.], 10, 104. Muthmann and Weiss, Annalen. 1901. 331, 1 (2Nd 1.70) <sup>3</sup> Popovier, Ber. 1908, 41.634 <sup>4</sup> Matignon, Compt. read., 1901. 133, 289, 1905. 140, 1637. Ann Chem. Phus. 1900, [viii.], 8, 243. Bourion, ibid., 1910, [viii.], 21, 19. of R. J. Mayer and Koss, Ber., 1902, 35, 2622. VOL. IV.

From aqueous solution the hexahydrate, NdCl3 6H2O, crystallises out in roso-coloured, monochine crystals of density 2 282 at 165°. The hevaly drate melts at 124° One hundred parts of water dissolve 987 parts at 13°, 1400 parts at 100° of the anhydrous chloride, the solution being in equilibrium with the hexahydrate. The concentrated aqueous solution dissolves neodymnum oxalate on warming, and a neodymnum oxalochlorule crystallises out when the solution is cooled.

The monohydrate, AdCl<sub>8</sub> H<sub>2</sub>O, is also known (p. 254)

The anhydrous chloride forms the following compounds with ammonia, the dissociation temperature being given with each formula -1

```
\begin{array}{ccc} {\rm NdCl_1\,8NH_3} & 79^{\circ} \\ {\rm NdCl_3\,11NH_3} & 26^{\circ} \\ {\rm NdCl_3\,12NH_3} & -10^{\circ} \end{array}
  360° C.
  NdCl<sub>3</sub> NH<sub>3</sub>
NdCl<sub>3</sub>2NH<sub>4</sub> 255'
NdCl<sub>3</sub>4NH<sub>4</sub> 157'
NdCl<sub>3</sub>5NH<sub>4</sub> 117°
  255,
```

Neodymium oxychloride, NdOCl, has been prepared (p. 255) Neodymium bromide, NdBi, resembles the chloride See p 255 Neodymium iodide, Ndl, also resembles the chloride Sec p 255

Neodymium bromate, Nd(BrO), 9H<sub>2</sub>O, forms pink hexagonal prisms. At 100° the dihydrate, Nd(BrO,) 2H2O, and at 150° the anhydrous salt is obtained, at higher temperatures decomposition takes place. See p. 256.

# NEODYMILM AND THE OXYGEN GROUP

Neodymium sesquioxide, or neodymia, Nd<sub>2</sub>O<sub>3</sub>, may be prepared by igniting the hydroxide, carbonate, intrate, oxalate, etc.—The oxide obtained by iginting the oxalate strongly has a pure sky blue colour. It dissolves readily in acids

As regards the existence of higher oxides of neodymium, the evidence is conflicting. Aner von Welsbach, Jones, and R. J. Meyer and Koss deny that a higher oxide may be prepared in the dry way, but Marc maintains that neodymia may be peroxidised in the presence of cerumi and praseodymium. Waegner also allirms the existence of a higher oxide,  ${\rm Nd}_1{\rm O}_7$ , and even describes its reflection spectrum, but Joye and Garmer have shown that this spectrum is due to a hydrated exide,  $Nd_2O_3H_2O$  Brauner states (without any details) that the exides  $Nd_2O_4$  and  $Nd_2O_6$  both exist. By the latter Brauner presumably means a hydrated peroxide,  $Nd_{Q_0}$  all  $Q_0$  or  $(OH)_Nd^{(0)}$  of  $Q_0$  corresponding to the known lauthanim and prascodynnum peroxides. The same authority also mentions the existence of an acetate of the constitution  $(\text{CH}_3(\text{CO})_2\text{Nd}^{111}(\text{O.O.O.O.Nd}^{111}(\text{OH})(\text{CO.CH}_3) + \text{H}_2\text{O}$ , analogous to the prascodynnum compound.2

Neodymium hydroxide. Nd(OH)<sub>30</sub> is obtained as a blue precipitate when an alkah hydroxide is added in excess to a solution of a neodymnum salt. When dired at 300° it leaves the hydrate 2Nd<sub>2</sub>O<sub>4</sub>3H<sub>2</sub>O<sub>5</sub> this, when heated to 525°, is converted into the hydrate Nd<sub>2</sub>O<sub>3</sub> H<sub>2</sub>O; and the latter at a bright red heat is converted into neodymia 3

<sup>&</sup>lt;sup>1</sup> Matignon and Trainoy, Compt rend., 1906, 142, 1042.

<sup>2</sup> Jones, Amer Chem J., 1898, 20, 315, Meyer and Koss, Ber., 1902, 35, 3710, Marc, Ber., 1902, 35, 2370, Wargner, Zeitsch amorg Chem., 1904, 42, 118, Joye and Garmer, Compt rend., 1912, 154, 510, Joye, Arch See phys. nat., 1913, [iv], 36, 41; Brauner, Proc. Chem. Noc., 1888, 14, 72, 1901, 17, 56, Zeitsch, anory, Chem., 1902, 32, 1

<sup>3</sup> Joye and Garmer, Compt rend., 1912, 154, 510, Joye, Arch. See phys. nat., 1913, [iv], 36, 41.

Neodymium sulphide, Nd2S3, the preparation and properties of which have been described (p. 259), is an olivegreen powder of density 5 179 at 11° (Muthmann and Stutzel)

Neodymium sulphate, Nd2(SO1) - For preparation and general properties, see p. 260. The only hydrate known as the monochine octahydrate, Nd  $(80_4)_4\,81l_20$ , the solubility of which, in grains of anhydrous sulphate per 100 grams of water, is as follows

For acid and basic sulphates, Nd(HSO<sub>1</sub>), and Nd(O<sub>1</sub>SO<sub>1</sub> respectively, see p 263 Acodymium casaum sulpletts, Nd (80<sub>11</sub>, Cs\_80<sub>4</sub> 3H O, norms lavender-blue crystals <sup>1</sup>

Neodymium molybdate, Nd,(NoO<sub>4</sub>) = Sec. p. 265 Neodymium tungstate, Nd,(NO<sub>4</sub>)  $_{\rm p}$  and complex time tates have been mentioned (p. 265)

Neodymium silicotungstate, Nd<sub>4</sub>(W<sub>4</sub>,SiO<sub>10</sub>)<sub>4</sub> - See p. 206 Neodymium uranate, Nd<sub>2</sub>(U<sub>4</sub>O<sub>10</sub>)<sub>4</sub> 18H O, is obtained as a yellow, crystalline solid by heating neodymium hydrocide with aqueous unauyl acetale 5

### NEODYMEM AND THE NURSERN GLOUP

Neodymium intride, NdN, has been described (p. 267) Neodymium intrate, Nd(NO), 6H2O A valuated solution of this salt in water at 25 contains 152 9 grains of analydrous intrate per 100 grains of water 6. Neodymum intrate is isodimorphous with bismuth ratiote (see p 234). The double salts Nd(NO), 2(NH), NO, 4H, Or Nd(NO), 2RbNO, III,0,5 and the double salts of the type 2Nd(NO<sub>2</sub>), 3M(NO<sub>2</sub>), 24H O are known (see p. 268)

# NEODYMEN AND THE CARBON GROUP.

Neodymium carbide, NdC, has been described (p. 270)

Neodymium carbonate, Nd, (CO, p., 411,O, has not been analysed The following double salts are known -9

$$\begin{array}{lll} {\rm Nd}_2({\rm CO}_3)_3|{\rm K}_2({\rm CO}_1|2H,0) & 2{\rm Nd}_2({\rm CO}_4)_3|{\rm SA}_2({\rm CO}_1|22H,0) \\ {\rm Nd}_2({\rm CO}_4)_4|({\rm NH}_4)_4({\rm CO}_4|H,0) & 2{\rm Nd}_2({\rm CO}_4)_4|{\rm SA}_4({\rm CO}_4)_4|{\rm Nd}_4({\rm CO}_4)_4 & 2{\rm Nd}_4({\rm CO}_4)_4|{\rm Nd}_4({\rm CO}_4)_$$

Neodymium ethylsulphate, Nd(CH,SO<sub>4</sub>), 9H\_O, has been described (p. 278)

Neodymium acetylacetonate, Nd(CH gCO CH CO CH ), has also been described (p. 279)

Neodymium oxalate, Nd, (C,O<sub>4</sub>), 1011,O<sub>5</sub> is obtained by double decomposition as a pale rose-red crystalline powder. For its solubility in water and

C Matignon (Ann. Chem. Theo., 1907, [vni.] 10, 110) mentions the existence of a pentahydrite, Ad(SO<sub>1</sub>, 5H<sub>2</sub>O<sub>2</sub>, bat does not describe its preparation.

Extrus, Zeitsch Krait Man., 1904, 34, 307. Dufer, Chem. Ze. tr., 1902, 4, 452.

Muthinaon and Rolig. Ber., 1898, 31–1718.

Biskerville and Holland. J. Limit. Chem. Soc., 1904, 26, 71.

Orloll, Chem. Zeit., 1907, 31, 1119.

James and Robinson, J. Limit. Chem. Soc., 1913, 35, 754.

Auer von Welsbach, Monatsk., 188. 6, 477.

Jantsch and Wigdorow, Zeitsch. among. Chem., 1911, 69, 221.

R. J. Meyer, Zeitsch. anong. Chem., 1904, 41, 97., heep. 271.

acids, see p 273 According to James and Robinson, the only stable hydrate m contact with water or aqueous needymum intrate at 25 has the composition  $\mathrm{Nd}_2(\mathbb{C}_2\mathbb{Q}_4)_3$  11H,O. The oxalate dissolves perceptibly in aqueous neodymnum initiate, and under certain conditions neodymnum oralonitrate  $\operatorname{Nd}_2(C_2O_1)$ ,  $\operatorname{1Nd}(\operatorname{NO}_1)$ ,  $\operatorname{1H}_2O$  or perhaps  $\operatorname{Nd}_2(C_2O_1)$ ,  $\operatorname{5Nd}(\operatorname{NO}_1)$ ,  $\operatorname{1H}_2O$ , is produced. The solubility curve for a temperature of 25' is shown in fig. 10The oxidate also dissolves in an aqueous solution of the chloride, and a neodymunii oralochloride exists

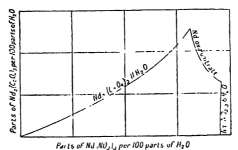


Fig. 40 — The system Nd  $(C_2O_4)_{\downarrow}$  -Nd(NO ) —H2O at 25 Solubility drigiam

Neodymium salts of organic acids.-- The following salts have been described

Neodymium acetate, malonate,  $^7$  sebacate,  $^4$  glycollate,  $^4$  cacodytate,  $^4$  dimethylphosphate," methyl, ethyl, propyl, and cobutyl sulphonates, methonetri-sulphonates, ethanedisulphonates, quenates, anisates, ocumbates, phonocyaectates, m-netroben enesulphonate, p dibromoben enesulphonate, a p dichloroben enesulphonate, 11 1 2 bromonitroben enesulphonate, 1 3 1 metasylenesulphonate,3 m-sulpholo a oute, camphorsulphonate, 8 hydroxynaphthalene 1-sulphonate?

### SAMARIUM

Symbol, Sm 12 Atomic weight 150 4 (O-16).

Samarum is one of the less common of the rare earth elements, being on the one hand much less abundant than the other elements of the cerum group, and on the other hand much more abundant than europaum, terbann, and thulium - It is about as plentiful as gadolinium

<sup>1</sup> James and Robinson J. Amer. Chem. Soc., 1913, 35, 751.
2 Pract and Lames, J. Amer. Chem. Soc., 1911, 33, 1830.
1 James, Hoben, and Robinson, that, 1912, 34, 27b.
4 Whittenior and James, that, 1913, 35, 127
6 Katz and James, that, 1913, 35, 872
6 Morgan and James, that, 1911, 36, 10
7 Endmann and Writh, Annah n, 1908, 361, 190
7 Holmberg, Zetten among Chem., 1907, 53, 83
9 Jantsch and Grunkrant that, 1913, 79, 305
10 Armstrong and Rodd, Proc. Roy. Soc., 1912, A. 87, 201
11 Rodd, that, 1912, A. 89, 202
12 Two symbols are in common use for samanum, viz., Sin and Sa. In this book the former has been adopted, since it is the symbol that was assigned to the element by its discoverer, Lecoq de Borsbaudian.

The occurrence, history, preparation, atomic weight, and homogeneity of samarnin have been already discussed in Chapters X, and XI

Another method for the separation of samarum from Lunthanum, prascodymium, and needymium may conveniently be mentioned here, its theoretical value has been pointed out by Matignon and Cazes, but its practical value does not appear to have been determined. The mixed oxides are converted into the mixed anhydrous chlorides by the method of Bourion or some other suitable process (p. 252), and heated to reduces in a stream or dry hydrogen The samarium is thus converted into the lower chloride SmCL, which may be isolated from the mixture by extracting the other chlorides with alcohol If, on the other hand, the mixed chlorides are dissolved in a large quantity of water, about one third of the samarium separates as hydroxide and the wished precipitate may be freed from small quantities of oxychlorides by dis-olving it in cold, dilute hydrochloric acid, in which the oxychlorides are insoluble.

Samarium 'is a vellow metal of density 7.7-7.8. It is as hard as steel and melts at 1300°=1400°. The metal quickly tarmshes in the an

#### COMPOUNDS OF SAMARIUM

The majority of the known compounds of samai ium are salts derived from the basic oxide  $\mathrm{Sm}[O_3]$  . They are pale yellow in colour and the absorption spectra of their solutions exhibit characteristic bands in the blue and violet. Two compounds of the type  $SmN_n$  however, are known, viz, the chloride and inclide

The equivalent conductivities (X) of samarium chloride and sulphate solutions at 25°C indicate that the salts are derived from a tank strong triacid base (v -litres per gram-equivalent) =  $^{1}$ 

```
32.59
                          -65 [8 | 130 36 | 260 72 | 521 14
   10129
                         1125
                                  120.8
           \lambda = 107.2
  126.1 - 131.3
  135.9
\mathrm{SR}_{\mathrm{G}}(\mathrm{SO}_1)_{+}v
               31 35
                         62.70 | 125.40 | 250.80 | 501.60
  10032
                31-17
                          -13/19
                                   51.59
  61.39
   72.76
   89 24
```

The compounds of samarum have been described mainly by Cleve, whose samaria was contaminated with a little emopia 1

Thermochemistry of Samarium. The following results have been obtained -

```
 \begin{split} & \left[ Sm_2O_3 \right] + 6HClar \\ & \left[ Sm_2O_4 \right] + 6(HCl) \\ & \left[ Sm_2O_4 \right] + 3[H,SO_4] \\ & \left[ Sm_2O_4 \right] + 3[H,SO_4] \\ & \left[ Sm_2Cl_4 \right] + 3[H,SO_4] \\ & \left[ 
  + 9163 Cds
  +128.1
  +1138
  1 37 1
   ٠,
  [\operatorname{Sm}_i(\operatorname{SO}_i)_i] + \operatorname{Aq}
  Sm (50) aq
  1 38 8
and
  \begin{split} & \left[ SmCL_{9}NH_{3} \right] + + 20.7 \text{ Cals} \;, \\ & \left[ SmCL_{9}2NH_{3} \right] + + 37.1 \;, \\ & \left[ SmCL_{9}3NH_{3} \right] + + 52.2 \;, \\ & \left[ SmCL_{9}4NH_{3} \right] = + 65.9 \;, \end{split}
  [SmCl_s5NH_s]
  4 780 Cds
  SmCL,8NH,
SmCL,95NH
  4-1116 ..
  \pm 126.6
   [SmCl<sub>2</sub>H 5NH] +1450 ",
```

Matignon and Cazes, Ann. Chem. Phys., 1996, [vm.], 10, 424
 Muthmann and Werss. An index, 1994, 331, 1., see pp. 229–250
 Aufricht, Innumeral Description (Berlin, 1994). Heyaweller, Zensch. physical d.

Chem., 1915, 89, 281.

4 Cleve, Chem. Acres, 1880, 53, 30, etc., see also Cleve, Cound. rend., 1883, 97, 91.

4 Cleve, Chem. Acres, 1880, 53, 30, etc., see also Cleve, Cound. rend., 1883, 97, 91.

Bull Son chim., 1885, [n.] 43, 162. Chem. Acres, 1883, 48. 9, 74. 1885, 51. 115;

Trans Chem. Son., 1883, 43, 362. Center Stensko Fet. Alsol. Fechandl., 1883, 40, 7.

Matignon, Compl. rend., 1906, 142, 276., Ann. Open. Phys., 1906, [vm.], 8, 402.

#### SAMARIUM AND THE FIVORINE GROUP.

Samarium fluoride, SmF<sub>3</sub> H<sub>2</sub>O, is obtained by double decomposition (('leve)

Samarium chloride, SmCl, is a very pale yellow, hygroscopic, crystalline solid which darkens when heated and melts to a brown liquid. For its preparation and properties, see p 252

From aqueous solution the hexahydrate, SmCl 611,0, separates in large, yellow tablets of density 2 383 (Cleve). The monohydrate, SmCl\_H<sub>2</sub>O<sub>2</sub> is also known (p. 254) — Samarrum aur who rde, SmCl, AnCl, 10H,0, crystallises in orange prisms, the chloroplatinate has the formula 2(SinCl<sub>3</sub> PtCl<sub>4</sub>)2111.0

Anhydrons samarnum chloride absorbs ammonia. The following compounds are known, their dissociation pressures being 760 mm, at the tempera tures given opposite the formulæ .-

| SmCl_XII                | 375° C | $\mathrm{SmCL}, 5\mathrm{NH}_3$       | 105' C |
|-------------------------|--------|---------------------------------------|--------|
| SmC1, 2N11,             | 210    | $SmCL_{i}SNH_{i}$                     | 76°    |
| $\mathrm{SmCl}[[3N11]]$ | 200,   | $\operatorname{SmCl}[9.5\mathrm{NH}]$ | 10,    |
| SmCL 4NH                | 155    | SmCT 11 5NH                           | 15°    |

Curiously enough, a compound with 6NH, does not appear to exist

Samarium oxychloride, SmO(L, 18 known (p. 255).
Samarium subchloride, SmCL, 18 prepared by strongly heating samarium chloride in a cirrent of pine, dry hydrogen or ammonia. It is a dark brown, crystalline solid of density 3-69 at 227, insoluble in alcehol, pyridine, carbon disalphide, chlorotorin, or benzene. It dissolves readily in water, giving a dark reddish-brown solution. The coloni slowly disappears, hydrogen being evolved and samarini hydroxide precipitated. The rapidity with which the colour disappears is greatly enhanced by shaking the solution ın aır .-- 1

$$68mCl_2 + 6H_2O = 48mCl_1 + 28m(OH)_1 + 3H_2$$
  
 $128mCl_2 + 6H_2O + 3O_3 + 88mCl_1 + 48m(OH)_1$ 

Samarium bromide, SmBr Seep 255 Thehevaliydrate, SmBr 6H<sub>2</sub>O, separates from aqueous solution in prismatic crystals of density 2 97, and forms an auribromide, SmBr , AnBr , 10H , 0 (Cleve)

Samarium todide, Sml, has been prepared by Matignon (p. 255)

Samarium subiodide, Sml, has been prepared by reducing the heated tri-rodide in a current of dry hydrogen  $^3$ 

Samarium bromate, Sm(BrO<sub>3</sub>) 911,0 - See p 256 This salt crystallises in yellow, hexagonal prisms, is converted into the dihydrate, Sm(BrO<sub>3</sub>), 2H<sub>2</sub>O<sub>5</sub> at 100°, the anhydrous salt at 150°, and decomposes at higher temperatures

Samarium iodate, Sm(103), 6H2O, forms a volummous, white, amorphous precipitate (Cleve)

Samarium periodate, SmIO, 411,0, may be obtained as an amorphous precipitate which becomes crystalline on standing (Cleve)

<sup>1</sup> Matignon, Compl. rend , 1902, 134, 1308 , Ann. Chim. Phys., 1 06, [vm.]. 8, 403 ,

Bounton, that, 1910, [viii ], 21, 49

2 Mattgnon and Tiannoy, Compt. vin.l., 1905. 140, 111. Jan Chim Phys., 1906, [viii ],

<sup>8, 407</sup> Matignon and Cares, Compt. rend , 1906, 142, 83 . Ann. Chim Phys , 1906, [viii.], 8, 417

### SAMARIUM AND THE OXYGEN GROUP.

Samarium sesquioxide, or samaria, Sm.O., may be prepared by the ignition of the hydroxide, carbonate, intrate, oxalate, etc. It is an almost white powder with a faint yellow tinge, and its density is 8 317 - It dissolves readily in acids. The corresponding hydroxide is a gelatinous precipitate that absorbs carbon dioxide from the air. The addition of ammonia al hydrogen peroxide to a solution of a samarium salt causes the precipitation of a hydrated percende,  $\mathrm{Sm_1O_9}$  eH2O or probably  $\mathrm{Sm_2O_7}\mathrm{eH_2O^{-1}}$ 

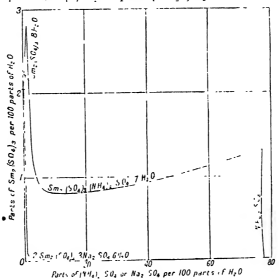


Fig. 41 — The system, Sm. So. , — Na SO. + H O and Sm. SO. + (NH. ]SO. – H<sub>2</sub>O at 2 $\sigma$  .— Solubility drag in

Samarium sulphide, Sm<sub>2</sub>S<sub>3</sub>, is a yellow solid of density 3.70<sup>2</sup>

Samarium sulphite, Sm2(SO.) 3H2O, has been obtained by Cleve as a

white, amorphous powder (see p. 260)

Samarium sulphate, Sm<sub>2</sub>(8O<sub>4</sub>) —For preparation and general properties, see p. 260. The monoclime bottahydrate, Sm<sub>2</sub>(8O<sub>4</sub>) SH<sub>2</sub>O<sub>5</sub> is the only hydrate known. The acid sulphate, Sm(HSO<sub>3</sub>), and the basic sulphate, Sm<sub>2</sub>O<sub>4</sub>SO<sub>5</sub> are known (p. 263). The following double sulphates have been the subset by Chem. described by Cleve ---

$$\begin{array}{lll} 2\mathrm{Sm}_2(\mathrm{SO}_4), 9\mathrm{K}_2(\mathrm{SO}_4) 3\mathrm{H}_3(\mathrm{O}_4) & \mathrm{Sm}_2(\mathrm{SO}_4), (\mathrm{NH}_4(\mathrm{SO}_4) \mathrm{SH}_2(\mathrm{O}_4)) \\ -\mathrm{Sm}_2(\mathrm{SO}_4), \mathrm{Na}_2(\mathrm{SO}_4) 2\mathrm{H}_3(\mathrm{O}_4) & \mathrm{Sm}_2(\mathrm{SO}_4), (\mathrm{NH}_4(\mathrm{SO}_4) \mathrm{SH}_4(\mathrm{O}_4)) \end{array}$$

Cleve, Bull. Soc. chim., 1885. [ii.1], 43, 53
 Edmann and Witth. Ann den. 1908, 361, 190., et. Morssan, Compt. rend., 1906, 131, 42, 850, 950

<sup>924.</sup> See p 259.
1 Fork, Zeitsch Kryst Min , 1900, 32, 250 , Dufet, Chem. Zente , 1902, 1 452. See p. 261

According to Keyes and James,1 the only double sodium and ammonium sulphates that exist at 25' have the formula 28m (SO<sub>4</sub>), 3Na,8O<sub>4</sub>6H,O and Sm<sub>2</sub>(SO<sub>1</sub>)<sub>3</sub> (NH<sub>2</sub>)<sub>2</sub>SO<sub>1</sub>7H<sub>2</sub>O. The results of their solubility measurements are shown graphically in fig. 41

Samarium selenite. - Sodium selenite gives with samarium sulphate a precipitate of a basic sclenite Acid samar rum sclenite, Sin (SeO.) H SeO. 411,0, is precipitated when aqueous solutions of samarium acctate and selemous acid are mixed (Cleve)

Samarium selenate, Sm<sub>2</sub>(SeO<sub>4</sub>)<sub>3</sub>. The octahydrate, Sm<sub>2</sub>(SeO<sub>4</sub>)<sub>3</sub> 8H<sub>2</sub>O<sub>4</sub> crystallises from aqueous solution at ordinary temperatures | Below 10, the dodecahydrate, Sm (SeO<sub>1</sub>), 1211,O, separates in shining, prismatic crystals The following double salts are readily soluble in water (Cleve) -

$$Sm_{2}(SeO_{4})_{3} \ K_{2}SeO_{4} \ 6H_{2}O \\ Sm_{2}(SeO_{4})_{3} \ (NH_{4})_{2}SeO_{4} \ 6H_{3}O$$

Samarium chromate. Potassium dichromate gives no precipitate with samarium intrate, but potassium chromate gives a vellow, crystalline pre cipitate of samarium potassium chromate,  $\mathrm{Sm}_{0}(\mathrm{CrO}_{4})_{3}$   $\mathrm{K}_{0}\mathrm{CrO}_{4}$  6H O (Cleve)

Samarium molybdate, Sm<sub>2</sub>(MoO<sub>1</sub>), forms small, lustrons crystals of density approximately 5 95. The double salt, SmNa(MoO<sub>4</sub>), has also been made (Cleve)

Samarium tungstate.—The metatungstate, Sm  $(W_1O_{1,i})$ , 35H O, forms readily soluble, topaz-vellow crystals of density 3 991  $-\Lambda$  section samarium tungstate is also known (see p. 265)

Samarium silicotungstate, Sm<sub>1</sub>(W<sub>11</sub>SiO<sub>10</sub>) —See p. 266.

# SAMARIUM AND THE NITROGEN GROUP

Samarium nitride. See p 267

Samarium nitrate, Sm(NO), 611,0, crystallises in extremely coluble, yellow needles, melting at 78-79 (Cleve). The double salts are described on p.  $268^{\circ 2}$ 

Samarium orthophosphate, SmPO, is obtained in the crystalline form by fusing sodium metaphosphate with samaria, allowing to cool, and extracting the mass with water. It is a crystalline powder of density 5.83 at 175°, msoluble in water (Cleve)

Samarium hydrogen pyrophosphate, 28mHP,0,7H,0, may be prepared by adding samarinia hydroxide to aqueous pyrophosphoric acid

(Cleve) The compound Sm<sub>2</sub>O<sub>3</sub> 5P<sub>2</sub>O<sub>5</sub> is also known

Samarium vanadates. Two hydrates of the salt Sm<sub>2</sub>O<sub>5</sub> 5V<sub>2</sub>O<sub>5</sub> have been prepared by Cleve, namely, red crystals of Sm<sub>2</sub>O<sub>1</sub>5V<sub>2</sub>O<sub>1</sub>28H<sub>2</sub>O and yellow crystals of Sm<sub>2</sub>O<sub>3</sub> 5V<sub>2</sub>O<sub>5</sub> 21H<sub>2</sub>O

# SAMARIUM AND THE CARBON GROUP

Samarium carbide, SmC<sub>2</sub>, has been described (p. 270) Samarium carbonate, Sm<sub>2</sub>(CO<sub>2</sub>)<sub>3</sub> 3H<sub>2</sub>O, loses 1H<sub>2</sub>O at 100° The following double salts are known (Cleve) -

> $\begin{array}{l} {\rm Sm}_2({\rm CO}_3)_3 \; {\rm K}_2{\rm CO}_5 \; 12 {\rm H}_2{\rm O} \\ {\rm Sm}_2({\rm CO}_3)_3 \; {\rm Na}_2{\rm CO}_3 \; 16 {\rm H}_2{\rm O} \end{array}$  $\operatorname{Sm}_2(\operatorname{CO}_3)_{\mathfrak{q}}(\operatorname{NH}_4)_{\mathfrak{q}}(\operatorname{CO}_3\operatorname{4H}_4O)$

Keyes and James, J. Amer. Chem. Soc., 1914, 36 634.
 See also Demarçay, Compt. rend., 1900 130, 1185

Samarium platinocyanide, 28m(CN), 3Pt(CN), 18H,O, crystallises in

yellow prisms having a blue reflex (rt. p. 272)

Samarium thiocyanate, Sui(CNS) 6H.O. forms prismatic crystals easily soluble in water. The double salt, Sm(CNS) [3HgrCN), 12H<sub>2</sub>O, is also known (Cleve)

Samarium ethylsulphate, Sm(C II SO<sub>1</sub>)<sub>3</sub> 9II O, has been described 278)

Samarium acetylacetonate,  $\mathrm{Sm}(\mathrm{CH}/\mathrm{CO}|\mathrm{CH}|\mathrm{OO}|\mathrm{CH}_{0})$  , has also been described (p. 279)

Samarium oxalate, Sm (C.O<sub>1</sub>), 10H O, is obtained as a vellow, crystalline precipitate by adding a solution of oxalic acid to one of a samarium salt For its solubility in water and acids, see p. 273. When potassium oxalate is used as precipitant, samarann polassann acadate, Sm.(C2O1), K.C.O1 5H.O. is obtained (Cleve)

Samarium salts of organic acids.—The following salts have been described

Samatum tormate, arctite, proposite, throacetate, algoritate, so social att, citrate, tartrate, malande, citratomate, schate, dimethylphosphate, cacodulate, thuladocallate, methylsulphomate thulsulphomate, propul sulphonate, sobut yest promate, nothanetresulphonate; ethanolesulphonate phonocravetate, m nitroben enesulphonate, p debeomoben enesulphonate, 10  $\frac{1}{1}$  | 4|  $\frac{2}{2}$   $\frac{1}{4}$   $\frac{1}{4}$   $\frac{3}{4}$   $\frac{1}{4}$   sulphomate, \ ^{4} \ and \ ^{8} \ hydrocenneighthalone \ ^{4} \ sulphomate)$ 

#### SAMARIPM AND BORON.

Samarium orthoborate, SmBO, is produced when samina is dissolved m molten boray at a white heat. It crystallises in very thin microcons se des, which may be extracted from the mass, after cooling, by carcial treatment with cold, dilute hydrochloric acid (Cleve)

Samarium metaborate,  $Sin(BO_2)_0$ , has been described (p. 282)

- Cleve Cherr No. 1883 53 30
   Pratt and Jones J. Lower Chem. 80 1911, 33 1 39
   James, Holien and Robinson. dod., 1912 34 276
   Whittemore and Limes, ib.d., 1913, 35 127
   Katz and Limes, ib.a., 1913 35 872
   Morgan and Tron. ibid. 1914, 36 10
   Erdmann and Wirth, Transler 1908 361 190
   Holmberg Zeelen away Chem. 1907 53 83
   Jantsch and Germlant. doc. 1914 79 505
   Armstrong red Rodd. Proc. Row. 866 1912 87, 204

### CHAPTER XIII.

# THE RARE EARTH ELEMENTS (continued).

### THE TERBIUM GROUP.

THE rare carth elements included in this group are only three in number, namely, europium, gadolinium, and terbium. Europium and terbium are rare in comparison with gardolimum, itself not one of the most abundant of the rare earth elements

By utilising bismuth as a separating element it is possible to effect a quantitative separation of the cerium and terbium groups by the crystallisation of the double magnesium intrates. At the same time a very good separation of europium and gadolimium is obtained. The most rapid method for eliminating terbium from gadolimum consists in fractionally crystallising the bromates. A summary of the various processes for the separation of the terbinin group is given in Chapter XI

Few compounds of europium and terbinin have been described.

# EUROPIUM.

Symbol, Eu. Atomic weight, 152 0 (O = 16).

Europuum ( $Z_{\rm c}, Z_{\rm 5}, Z_{\rm 5}$  and  $S_{\rm c}$ ) is one of the least abundant of the rare th elements. Monazite sand contains about 0.002 per cent of europia,  $^{1}$ Europium has been detected spectroscopically in the chromosphere of the sim and in the stars a-Bootis and B-Gemmorium?

Europium compounds are pale pink in colour. Europia was isolated by Demarçay, and prepared in a state of purity by Urbain and Lacombe in 1904. Subsequently it has been prepared by James and Robinson 3. The methods employed have been aheady described (p. 348).

Europium chloride, EuCl., has been prepared by Bonnon (p. 252) Europous chloride, EuCl., is prepared by heating the anhydrous trichloride to 100' 450' in a current of dry hydrogen. It is a colourless, amorphous solid, which dissolves in cold water, forming a neutral solution The aqueous solution oxidises on boiling - 1

 $12\text{EuCl}_{3} + 3O_{2} + 6H_{2}O = 8\text{EuCl}_{3} + 4\text{En(OH)}_{3}$ 

<sup>1</sup> Urbam and Lacombe, Compt. rend., 1904, 138, 627.

<sup>2</sup> Lunt, Pro. Ron. Soc., 1907, A, 79, 118.

<sup>3</sup> Demanday, Compt. rend., 1909, 130, 1019, 1469, 1901, 132, 1494, Urbam and Lacombe, thad., 1903, 137, 792, 1904, 138, 84, 627, J. Chim. phys., 1906, 4, 31, 105; James and Robinson, J. Amer. Chem. Soc., 1911, 33, 1393. See also Steddart and Hill, thid., 1911, 33, 1076

<sup>4</sup> Urbam and Bourion, Compt. 12nd., 1911, 153, 1155

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Europium sesquioxide, or europia, Eu<sub>2</sub>O<sub>5</sub>, produced by igniting the oxalate or sulphate, is a pale pink powder

Europium sulphate, Eu (80), is obtained by heating the octohydrate, Eu<sub>2</sub>(80<sub>4</sub>), 8H<sub>2</sub>0, to 375. It is completely decomposed at 1600.

Europium ethylsulphate, Eu(C,H,SO<sub>1</sub>) 9H<sub>2</sub>O, has been described

Other organic salts.- The tormate,2 quinate, pyrominente, minitrabenzenesulphonate,2 and 1 4 2 bromontroben encsulphonate3 have been

#### GADOLINIUM

Symbol, Gd. Atomic weight, 157-3 (O +16)

Gadolimin (Ya and victorium) is, like samarium, norther one of the most nor one of the least abandont of the rare earth elements. It forms a colourless oxide and a series of colourless salts, which exhibit no selective absorption in the visible region of the spectrum, but show an ultraviolet absorption spectrum. Gadolinia was doubtless one of the main constituents of the "terha" obtained by the carbor workers in the field of rare earth chemistry

Since gadolimim was discovered by Mariginae and examined by Leeing de Borsbandran, the isolation of pure gadolina has been studied by Bettendorf, Benedicks, Demarcay, Marc, and by Urbani and Lacombe 1

The salts of garlolimium have been described in unity by Benedicks 5

Gadolinium fluoride, GdF, prepared from appeous gadotinium sulphate and hydrofluorie acid, is a coloniless, gelatinous precipitate which becomes gramdar when warmed "

Gadolinium chloride, GdCl, is a white, hygroscopic crystalline is as (p. 252). It forms a heyabydrate, GdCl, 6H,O, of density 2/12 - Gadolinium anichlaride, GdCl, Ant.1 10H.0, crystallisis in yellow tables of specific gravity 2.71. Gadelinium platinichloride, GdCl, PtCl<sub>1</sub> 10H<sub>2</sub>O, torms orangered crystals of density 2.72

Gadolinium oxychloride, GdOCl, has been prepared (p. 255)

Gadolinium bromide, GilBi , is known (p. 256), and also the hexahydrate, GdBr, 6H<sub>2</sub>O<sub>5</sub> which crystallises in rhombie tables of density 2.84

Gadolinium oxide, or gadolinia, Gd<sub>2</sub>O<sub>3</sub> is a white powder of density 7/107 at 15%. It is hygroscopie, absorbs earbou dioxide, and is readily soluble in acids. The gelatinous historiale, Gh(OH), also also be carbon dioxide rapidly

Gadolinium sulphide, Gd S is a hygroscopic, yellow solid or density It is slowly decomposed by water, quickly by acids?

Gadolinium sulphate, Gd2(SO1)3 - The octahydrate, Gd2(SO4), SH2O, is

<sup>7</sup> Endmann and With, Jano'en 1908 361, 190

the only hydrate known, and forms monoclinic crystals (p. 261). The solubility (in parts of anhydrous salt per 100 of water) is as follows ---

| 0°   | 10°  | 14~  | 25°  | 314  |
|------|------|------|------|------|
| 3.98 | 3 33 | 2.80 | 2.40 | 2.26 |

The solubility in a saturated solution of sodium sulphate is such that at 25°C the solution contains 0.6 gram of anhydrons gadolimum sulphate per htre, gadolinium sodium sulphale, Gd  $(80_1)$ , Na  $(80_1)$ , Na  $(80_1)$ 211,0, being the stable solid phase  $^4$ . The corresponding potassium salt is also known (Benedicks)

Gadolinium acid selenite, Gd<sub>2</sub>(SeO<sub>3</sub>)<sub>3</sub> H<sub>2</sub>SeO<sub>3</sub> 6H<sub>2</sub>O<sub>3</sub> obtained from gadolimum acetate and seleme acid, is an amorphous precipitate which becomes crystalline on standing

Gadolinium selenate, Gd (SeO), and the octa- and deca hydrates are known (p 264) The octahydrate separates from aqueous solution at 100', the decallydrate at the ordinary temperature. The double salt, quidolineum potassium sclenate, Gd<sub>2</sub>(SeO<sub>1</sub>), 3K<sub>2</sub>SeO<sub>1</sub> 4H<sub>2</sub>O<sub>5</sub> is also known

Gadolinium silicotungstate, Gd<sub>1</sub>(W<sub>1</sub>/SiO<sub>10</sub>), See p. 266

Gadolinium nitrate, Gd(NO<sub>2</sub>), 611.0, separates from aqueous solution in large, triclinic crystals of density 2/33, and melts at 91°2. From concentrated intric acid the pentahydrate, Gd(NO.) 5H<sub>2</sub>O, crystallises out in prisms of density 2.41 Of all the rare earth intrates, this salt is the least soluble in intric acid. It melts at 92%

Gadolinium ammonium uitrate,  $Gd(NO_3)/2(NH_4)NO/4H_5O_5$  is an extremely soluble, dehquescent salt For other double nitrates, see p. 269

Gadolinium vanadate, GdV<sub>10</sub> 13H<sub>2</sub>O, crystallises in vellow, triclime prisms of density 2 66

Gadolinium carbonate, Gd<sub>2</sub>(CO<sub>2</sub>), 1311,O(t), is obtained by passing a current of carbon dioxide for a long time through an aqueous suspension of gadolimium hydroxide - The basic carbonate, Gd(OH)CO H.O. is also known

Gadolinium platinocyanide, 2Gd(CN), 3Pt(CN), 21H<sub>2</sub>O, forms thombie crystals isomorphous with the yttrium salt

Gadolinium oxalate,  $\operatorname{Gd}_{*}(C_{1}O_{1})_{*}10\Pi_{*}O_{*}$  is obtained by precipitation as a white, incrocrystalline powder. It separates from concentrated in monoclinic crystals. For its solubility in sulphuric reid, see p. 273 The decalydrate loses 6H,O at 100

Gadolinium ethylsulphate, Gd(C<sub>2</sub>H<sub>1</sub>SO<sub>1</sub>), 9H<sub>2</sub>O = See p. 278.

Gadolinium acetate, Gd(C2H3O3)3 HI2O, forms soluble triclinic crystals of density 1 611 The proposate, dycollate, malonate, dimethylphasphate, 1 - 1 - 2 homonitroben zenesulphonate, minitroben enesulphonate, plubborobenzenesulphonate, and p-drbromoben zenesulphonate  $^{7.8}$  have also been described. Gadolinium metaborate,  $\mathrm{Gd}(\mathrm{BO}_2)$ , has been described (p. 282)

Bissel and James, J. Amer. Chem. Soc., 1916, 38, 873
 Deman, ay, low ett., von Lang and Hattinger, Annaden, 1907, 351, 150
 See p. 268.
 Beneche's, Zeitsche anna Chem., 1900, 22, 393.
 Holmberg, ibid. 1907, 53, 83
 Jantsch and Gronkraut, ibid., 1913, 79, 305
 Beneche's, Zeitsche and Gronkraut, ibid., 1918, 79, 305

Edbianin and Wirth, Anaden, 1998, 361, 190
 Armstrong and Rodd, Peor Roy, No., 1912, A, 87, 204.
 Rodd, that, 1913, A, 89, 292
 Katz and James, J. Inter Chem. Soc., 1913, 35, 872

<sup>10</sup> Morgan and James, chid , 1916. 36, 10.

# TERBIUM

Symbol, 1b. Atomic weight 159 2 (O 16)

Terbinia ( $Z_{s}, Z_{s}, \Gamma_{s}$  iomina, and incognition), notwithstanding the fact that its existence was forefold by Mosander in 1813, is one of the least abundant of the rare earth elements. Its curious history, and also the processes by which pane terbia may be isolited, have been already given in Chapters X and XI.—The preparation of pure terbia was first accomplished by  $U_{\mathrm{rb, m}}$ 

Terbia is white and the salts of terbium are colourless. The most interesting compound of terbium is the peroxide. Only a few terbium compounds have been described?

Terbium chloride, TbCl, prepared by Romion (p. 252), ery tallises in colonless needles which melt to a colonless liquid. The loxalivdrate, TbCl\_6H.O, forms hygroscopic, colourless, prismatic crystals

Terbium bromide, Tidh See je 255

Terbium sesquioxide, or terbia, Theo, is a white solid obtained by heating the peroxide in hydrogen.

Terbium peroxide, Tb<sub>1</sub>O<sub>0</sub> is produced by cilenting the terbium salts of volatile acids in an Prepared by the ignition of the oxalite in a muffle furnace, it is a dark brown powder, but when obtained from the sulphate at a white heat it is black. The strongly calcined peroxide is not attacked by cold innicial acids and is only slowly acted upon by hot, dilute acids

Owing to the ready formation of this percende and its intense coloning power, the presence of even traces of terbia in curopia, gadolinia, dysprosia, and vttma is easily detected

Terbium may be estimated quantitatively by igniting the oxalate and weighing the residual peroxide

Terbium sulphate, Tic(80), is prepared by dehydrating the octahydrate, Tb (80<sub>4</sub>) 8H O, at 360

Terbium silicotungstate, Th<sub>i</sub>(W<sub>12</sub>SiO<sub>10</sub>) See p. 266

Terbium nitrate, The NO ) 6HaO, forms colourless, monochine needles, m p 89 3

The propromate, pyromovate, and 1/4 Diromonitraten encodphonates have also been described

Urbein Cound (read), 1904-139, 7 br. 1905-141-521, 1909, 149, 37, J. Chim. phus.,
 1906-4, 531. See also Jaines and Berrl. J. June 1906, 149, 37, 1914, 36, 2009, and of
 Marc. Berr., 1902-35, 2382. Lett. Zeitsch. ann. g. Cherr., 1905, 43, 207. Policit., Chem. N. ns., 1905, 92, 3
 Urbain, Invest., and Compt. read., 1906, 142, 957. Urbain and Jaid ch. rbid., 1908, 436, 197.

<sup>3</sup> Grant and James, J. Iner Chem. Soc., 1915 37 2002

### CHAPTER XIV

# THE RARE EARTH ELEMENTS (ronelnded)

### THE YTTRIUM GROUP

The elements of this group are dysprosium, holimum, yttrium, erbinin, thulium, ytterbrum, luteerium, and celtium. All are rare in comparison with yttrum itself, ytterbum being perhaps the most abundant much less abundant than is commonly supposed.

The compounds of the elements of this group have received little attention at the hands of chemists. Attention has been directed mainly towards the problem of separating the actual earths of the group from one another, the results, however, still leave much to be desired, and considerable work still remains to be done in connection with the yttria earths

In the following brief account the elements are discussed in their serial order.

# DYSPROSIUM

Symbol, Dy Atomic weight, 162 5 (O -16)

Dysprosium  $(Z_0,\,Z_2,\,\Delta,\,$  and demonant  $^1)$  is one of the least abundant of the rare earth elements. Its salts were isolated in a pure state by Urban in 1906. They are yellow or greenish yellow in colour. Only a few dysprosium compounds have been described 2

Dysprosium chloride, DyCl, crystallises in pale vellow, pearly spangles, and melts to a coloniless liquid. It has been prepared by Bourion (p. 252) The chloride forms a hexahydrate, DyCl 611 O

Dysprosium oxychloride, DyOCl, is also known (p. 255)

Dypsrosium bromide, DyBr —See p. 255

Dysprosium bromate, Dy(BrO<sub>3</sub>)<sub>9</sub>.9H<sub>3</sub>O, forms shining, pale yellow, hexagonal needles, m.p. 78°. By prolonged heating at 110′, the trihydrate, Dy(BrO<sub>3</sub>)<sub>3</sub>.3H<sub>2</sub>O, is obtained <sup>3</sup>

Dysprosium sesquioxide, or dysprosia, Dy<sub>2</sub>O<sub>3</sub>, is a white, highly

paramagnetic solid (p. 256). No peroxide is known

Dysprosium sulphate, Dy (SO<sub>4</sub>),, may be obtained by dehydrating the octahydrate, Dy (SO<sub>4</sub>), at 360°. The latter forms brilliant yellow cig stals, stable at 110°.

 $<sup>^{1}</sup>$  Rowland's element demonsion was characterised by the spectrum line  $\lambda 1000$  6, which

<sup>-</sup> Advances element animatum was characterised by the spectrum the A1000 6, which belongs to dyprosum (see p. 308 - See Rowland Chem News, 1891, 70-68

<sup>2</sup> Urbam, Compt. rend., 1906, 142, 785. 1908, 146, 922, Urbam and Dementioux, 2bid, 1908, 143, 598. Urbam and Jantsch, 1bid., 1908, 146, 127, Jantsch and Ohl, Ber., 1911, 44, 1274

<sup>3</sup> Jantsch and Ohl, loc. ct

**Dysprosium selenate,** Dy/SeO<sub>4</sub>V, may be obtained by dehydrating the yellow, crystalline octahydrate, Dy/SeO<sub>4</sub>V, 8H<sub>2</sub>O<sub>5</sub> at 200°.

Dysprosium chromate, Dy (Crops 1011 O, obtained by double decomposition, is a greenish-yellow, microcrystidline powder. At 150° it loses 3.511,0, and above that temperature decomposition occurs saturated solution of the decally drate at 25° contains 10 grains of the

Dysprosium nitrate, Dy(NO<sub>2</sub>) 5H<sub>2</sub>O, melts at 88 6° and resembles the corresponding bismuth salt

Dysprosium phosphate, DyPO, 511,0, a gelatmous precipitate which slowly turns crystalline, is a pale yellow solid which may be dehydrated at 200° 1

Dysprosium carbonate, Dy (CO<sub>3</sub>), 1H<sub>9</sub>O<sub>5</sub> an insoluble powder formed by treating an aqueous suspension of dysprosium hydroxide with a current of carbon dioxide, loses 311,0 at ordinary temp ratures. In contact with saturated aqueous ammonium carbonate, it is slowly converted into sparingly soluble, crystalline dispression ammonium carbonate,  $(N\Pi_1)$  Dy $(CO_1)_3$   $\Pi_2O_5$ which begins to lose ammonia at 60°

Dysprosium oxalate, Dy  $(C,O_1)_1$  1011.0, is precipitated as microscopic prisms, insoluble in water. One litre of normal sulphuric acid at  $\pm 0^\circ$  dissolves 1893 grams of the oxalite (anhydrous). Disprosium polassium oxalate, KDv(C<sub>2</sub>O<sub>4</sub>), 3H<sub>2</sub>O<sub>5</sub> precipitated when solutions of dysprosium intrate and

potassuum oxalate are mixed, is reachly soluble in dilute acids!

Dysprosium platinocyanide, 2Dv(CN), 3Pt(CN), 2HLO, forms cinnabit ted crystals with a green tellex (et p 272). The tormate1 and acctate 1 are also known

### HOLMIUM.

Symbol, Ho Atomic weight, 1645 O 165

Holmmin is one of the least abundant of the rare earth elements. The isolation of pure holma has been a tack of great difficulty. The methods that have been used are described elsewhere (p. 356). The best holma yet obtained (1916) was extracted from the mineral envenite by Holmberg. It contained traces of erbia and dysprosia

Holmia, Ho2O, is a pale yellow, highly paramagnetic powder. The salts of holmium are yellow with an orange tinge

### YTTRIUM

Symbol, 1 Y Atomic weight, 88 7 (O 16)

Yttrium is the most abundant of the rare earth elements of the yttrium group Since, however, minerals rich in this group are much less abundant than minerals rich in the ceruin group, and the latter immerals are usually

<sup>1</sup> Jantsch and Ohl, loc. ed

<sup>2</sup> Sometimes called neoholmium the original "bohoum" being neoholmium plus

dyspresium Monthey Min Chem Min Ched , 1911, 4, No. 2 and 10 Zeasch award Chem , 1911, 77, 226
4 Two symbols are in use for yttimor, viz., Y and Y: The symbol Y used by

Berzelius and by all the early workers on the raic earths, has been adopted here.

very poor in yttroom, it is doubtful whether yttroom is as abundant as any of the cerum metals, samarium excepted

The occurrence, history, preparation, atomic weight, and homogeneity of

yttrium have been already discussed in Chapters X and XI

Yttrium. -The metal has not yet been prepared in quantity in the massive form. One or two chemists have prepared it as a grey, metallic powder, the chemical properties of which were similar to those of cerium and lanthanum, but they did not work with pure yttrium compounds.1

### COMPOUNDS OF YTTRIUM.

The compounds of yttrum have been described mainly by Cleve and Hoghind? Those derived from coloniless acids are themselves coloniless and exhibit no selective absorption

### YEIRIUM AND THE FLUORINE GROUP.

Yttrium fluoride, YF, See p. 252. The hemilydrate, 2YF, H<sub>2</sub>O, is obtained by double decomposition (Cleve)

Yttrium chloride, 'ACl, - For the preparation and properties of the anhydrous salt, see p 252. It is a hygroscopic, colourless, crystalline solid, perceptibly volitile at a bright red heat

The hexahydrate, YCl 6H O, crystallises in colourless, monoclinic prisms and melts at 156° to 160°. The monohydrate, YCl, H2O, is also known 251)

Ytterum anrichloride, VCI, 2AuCl 16H,O (Cleve), and attrium mercuric

chlorate, <sup>1</sup> YCl 3HgCl, 9H,O, are known Yttrium bromide, YBr<sub>9</sub>, crystallises from aqueous solution as the hydrate, YBi; 9H,O (Cleve)

Yttrium chlorate, Y(ClO), 9H2O, obtained by double decomposition between yttrium sulphate and barnun chlorate, crystallises from water in colourless needles (Cleve)

Yttrium perchlorate,  $Y(ClO_{1/3}, 91I_2O_3)$  is a very hygroscopic salt, soluble in water and alcohol (Cleve)

Yttrium bromate, Y(BrO<sub>2</sub>) 911.O, forms colourless, hexagonal prisms At 100' the trihydrate, Y(BrO) 3H2O, is produced (see p. 256)

Yttrium iodate, Y(IO<sub>3</sub>), 3H<sub>2</sub>O<sub>5</sub> is obtained from yttrium intrate and

iodic acid as a white precipitate (Cleve)

Yttrium per-iodate.—Two per iodates have been prepared by Cleve,  $\mathbf{Y_2O_3}$ ,  $\mathbf{I_2O_7}$  8  $\mathbf{II_2O}$  and  $\mathbf{3Y_2O}$ ,  $\mathbf{2I_2O_7}$  6  $\mathbf{II_2O}$ 

<sup>1</sup> For references, see pp. 229-230. S. Meyer (Monatch. 1899, 20, 793) gives the density of yttirum (powder) as 3 80 at 15°. Owen (Ann. Physik. 1912, [tv.], 37, 657) says that yttirum is paramagnetic. Pure yttira, however, is diamagnetic.

On the paramagnetic | Thie ytting, however, is diamagnetic |
 On the paramagnetic | Thie ytting alloys, see Siemens and Halske, D R.P. No. 116,503
 Cleve and Hegland, Bihama K. Srenska Vet -Akad Handl , 1873, 2. Bull Soc chem. 1873 [u.], 18-193, 289 | Ber., 1873, 6, 1467 | Cleve, Bihama K. Stenske Vet -Akad Handl , 1871, 2, No. 12 | Bull Soc chem., 1874, [u.], 21, 314 | ef Popp Angalen, 1864,

<sup>131, 197

\*\*</sup> Matignon, Compl. ren! 1902, 134, 1308, Ann Chim Phys., 1906, [viii.], 8, 433, Bourton, ibid., 1910, [viii.], 21, 49

\*\* Popp, Annelen, 1864, 131, 1/7.

# YITRIUM AND THE OXYGEN GROUP

m sesquioxide, or yttria, Y<sub>2</sub>O<sub>3</sub>, is a pine white powder of density 4.04 at 10 C<sup>4</sup> It is feebly diamagnetic (p. 257)

Yttria absorbs carbon dioxide from the air, liberates ammonia from

ammonium salts, and dissolves readily in acids

When dissolved in molten calcium chloride and cooled, yttma assumes the crystalline state, crystallising in trapezohedra.2 For further information concerning yttim, see p. 256

Yttrium hydroxide, Y(OII), is obtained as a gelatmous precipitate from solutions of yttrium salts by the addition of excess of a soluble hydroxide. It readily absorbs curbon droxide. When the precipitant is a mixture of ammonia and hydrogen peroxide, hydrated yttrium peroxide,

Y<sub>4</sub>O<sub>9</sub> xH<sub>4</sub>O or perhaps Y<sub>1</sub>O<sub>8</sub> rH<sub>4</sub>O<sub>1</sub> is produced <sup>3</sup>
Yttrium sulphide, Y<sub>2</sub>S<sub>2</sub> is a yellow solid which resembles the other rare earth sulphides <sup>3</sup> A green, crystalline, insoluble double sulphide,  $Y_2S_1$  Na<sub>2</sub>S<sub>1</sub> is obtained by heating a mixture of yttim and sodium chloride to  $1000^\circ$  in a stream of hydrogen sulphide, and washing the product with cold water 5

Yttrium sulphite,  $Y_2(SO_4)$ , 3H,0, has been prepared (p. 260) Yttrium sulphate,  $Y_2(SO_4)$  — For preparation and general properties, see p 260. The specific heat of the anhydrous sulphate between 0° and 100° is 0 1319 6

The monoclime? octahydrate, Y<sub>2</sub>(SO<sub>4</sub>)<sub>2</sub> SH<sub>2</sub>O<sub>5</sub> is the only hydrate known At 25' the saturated solution contains 5.38 grams of anhydrous sulphate per 100 grams of water 5. Its solubility diminishes with rise of temperature

For and and basic sulphates, see p. 263 The following double sulphates are known (Cleve) -

$$\begin{array}{lll} 2Y_2(8O_4)_+3K_2SO_4 & & & Y_2(8O_4)_+\lambda_4.8O_4\\ -Y_2(8O_4)_+4K_2SO_4 & & & Y_2(8O_4)_+(\lambda\Pi_4)_2SO_49\Pi_2O \end{array}$$

The addition of sodium sulphate increases the solubility of yttrium sulphate at first, and then causes the solubility to diminish rapidly diminution in solubility is associated a change in solid phase, from hydrated yttrum sulphate to double salt. Solutions supersaturated with respect to the double salt may remain in the metastable state for several months. The solubility curve for a temperature of 25' is shown in fig. 12.5

Yttrium dithionate, Y<sub>2</sub>(S<sub>2</sub>O<sub>6</sub>)<sub>3</sub> 18H<sub>2</sub>O, is a very soluble salt (Cleve)

<sup>&</sup>lt;sup>1</sup> R. J. Meyer and Whotmen, Zeitsch among Chem., 1913, 80, 7. Previous results are 5-03 (Cleve), 5-05 (Nilson and Pettersson), 1-51 (Muthmann and Bohm), and 5-32 to 5-38 (Tanatar and Volpansky, J. Ruse Phys. Chem. Soc., 1910, 42, 90). They suggest very strongly that the yttim used by Cleve, by Nilson and Pettersson, and by Tanatar and Volpansky area purpose. Stronger that the ettra used by Cleve, by Khison and Tenterson, and by Voljansky, was impure

2 Dubon, Compt. rend., 1888, 107, 99

3 Cleve, Bull. Soc. chim., 1885, [n], 43, 53

4 See p. 25<sup>n</sup>, also Cleve, Bull. Soc. chim., 1874, [n], 21, 344.

5 Dubon, Compt. rend., 1888, 107, 213

5 Nilson and Petterson, Compt. rend., 1880, 91, 232, Eer., 1880, 13, 1459

7 Kraus, Zeitsch. Kryst. Min., 1901, 34, 307, see p. 261.

8 James and Holden, J. Amer. Chem. Soc., 1913, 35, 559.

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Yttrium selenite, Y2(SeO3)3 12H2O, is obtained as a white precipitate by adding sodium selenite to a solution of yttrium sulphate, and is converted by warning with aqueous selemous acid into the acid selemite,  $Y_g(SeO_g)_gH_gSeO_gHI_gO$  (Cleve)

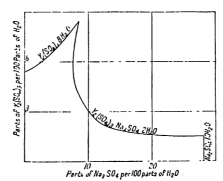


Fig. 12 —The system Y (804). Na<sub>2</sub>804—H<sub>2</sub>O at 25° C. Solubility diagram

Yttrium selenate,  $Y_2(\mathrm{SeO}_4)_3$ . The monoclinic octahydrate,  $Y_2(\mathrm{SeO}_4)_4$  8H<sub>2</sub>O<sub>4</sub> and the orthorhombic decahydrate,  $Y_2(\mathrm{SeO}_4)_4$ 10H<sub>2</sub>O<sub>5</sub> are known (see p. 261), also the following double sulphates --

$$Y_2(\operatorname{SeO}_1)_+ K_2 \operatorname{SeO}_1 \operatorname{6H}_2 O = Y_2(\operatorname{SeO}_1)_3 (\operatorname{NH}_1)_2 \operatorname{SeO}_1 \operatorname{6H}_2 O$$

which are readily soluble in water,

Yttrium tungstate,  $Y_i(W|Q_i)_i$  - See p. 265 Yttrium silicotungstate,  $Y_i(W|_{L_p}SiO_{10})_i$  - See p. 266

## YTIRIUM AND THE NITROGEN GROUP

**Yttrium nitrate,** Y(NO.)  $[6H_2O]$ , forms deliquescent, triclinic crystals <sup>1</sup> and loses  $[3H_2O]$  at [100]. The trimitrate, Y(NO.)  $[3H_2O]$ , crystallises from very concentrated intric acid." A dihydrate has also been described. When heated, yttrium intrate first forms basic intrate and subsequently at very high temperatures leaves a residue of yttria. Yttrium intrate is isodimorphous with bismuth intrate (see p. 234)

A saturated aqueous solution of the intrate at 25° has a specific gravity (25/25) of 17446 and contains 1416 grams of anhydrous intrate per 100 grams of water 4

The only basic yttrium intrate that exists at 25 has the composition 3YaO 4NaO 2011,O It is stable in air and in the presence of a solution of yttimm intrate containing one-fourth its weight or more of anhydrous

Von Lang and Haitinger, Anna'en 1907, 351, 450
 Demarçay, Compt. read., 1900, 130, 1019.
 Tanatai and Voljansky, J. Russ. Phys. Chem. Soc., 1910, 42, 586

<sup>4</sup> James and Pratt, Jf Amer. Chem. Sec., 1910, 32, 873.

intrate, and from such a solution it can be recrystallised (see fig. 43). It is not affected by absolute alcohol, but is decomposed by water  $^{\rm T}$ 

Yttrium orthophosphate, VPO, occurs in nature as the mineral senotime. An ammoniaeal solution of ammonium orthophosphite when added to vttrium intrate precipitates a white dilividrate, VPO<sub>1</sub>2H<sub>2</sub>O . The amorphous phosphate may be crystallised from molten yttrium chloride?

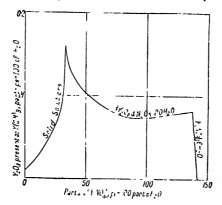


Fig. 13. –The system Y O —  $\Sigma_2O_3$  - H O at 2.7 C Solubility diagram

first ytterion orthopho photo, Y (HPO)), may be obtained as an amorphous precipitate by adding disodium orthophosphate to a solution of a yttrium salt (Cleve)

Yttrium hydrogen pyrophosphate, 2YHP,O, 7H,O, has been prepared by Cleve - The following compounds have also been described -

Yttrium metaphosphate, V(PO), is obtained by heating attria with molten metaphosphoric acid and removing excess of reid with water (Cleve)

YTEREM AND THE CARBON GROUP

Yttrium carbide, YC, has been described (p. 270) Yttrium carbonate, YaCO (311,0, and the following double salts are known (see p. 271)

$$Y_2(\mathrm{CO})_1 \, \mathrm{Na}_2 \, \mathrm{CO}_{-} \, \mathrm{HI}_2 \mathrm{O}_{-} = - Y_2(\mathrm{CO})_1 \, \mathrm{CNH}_2 \, \mathrm{CO}_{-} \, \mathrm{2H}_2 \mathrm{O}_{-}$$

Yttrium platinocyanide, 2Y(CN) 3Pt(CN), 2HLO torms and crystals having a green reflex (see p. 272).

James and Pratt, J. Amer. Chem. Soc., 1910, 32, 873
 Radominsky, Compl. rend., 1875, 80, 501
 Johnson, Ber., 1889, 22, 976., Wallroth, Bull. Soc. Chem., 1883, [ii], 39, 316.
 Duboin, Compl. rend., 1888, 107, 622.

Yttrium thiocyanate, Y(CNS),  $\Pi_2O$ , and likewise the double salt, Y(CNS),  $3\Pi_2(CN)$ ,  $12\Pi_2O$ , are known (see p. 272)
Yttrium oxalate,  $Y_2(C_2O_1)$ ,  $9\Pi_2O$ , is obtained by double decomposition as a white, crystalline precipitate. For its properties, see p. 273.

The only stable yttrium potassium oxalate at 25° has the formula  $Y_2(C_2O_4)$ ,  $3\Pi_2O_4$ ,  $3\Pi_2O_5$ , and is decomposed by water. Fig. 44 shows the solubility diagram, as determined by Pratt and James 1

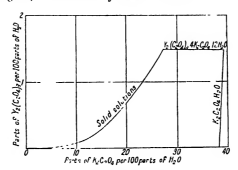


Fig. 14 —The system  $Y_2(C_2O_4)_3 + K_2C_2O_4$ -  $II_2O$  at 25° C. Solubility diagram.

Yttrium ethylsulphate, Y(C<sub>2</sub>H<sub>5</sub> SO<sub>4</sub>)<sub>3</sub> 9H<sub>2</sub>O, is known (p. 278). Yttrium acetylacetonate, Y(CH<sub>3</sub> CO CH CO CH<sub>4</sub>)<sub>5</sub> — See p. 279.

Other organic salts -The following salts are known -

Yttinum formate, 2, 3 acetate, 2, 8 propionate, 11 isobutyrate, 11 succinate, 2, 8, 4 tartrate, 2, 8, 10 glycollate, 5, 9 citrate, 10 lactate, 11 cacodylate, 5 schacate, 6 malonate, 8, 10 fumarate, 11 methylsulphonate's ethylsulphonate, methanidisulphonate, methanetrisulphonate, 5 crotomate, 11 malate, 11 citraconate, 11 benzoate, 11 phenylacetate, 5 phenogyacetate, 5 salicylate, 5 phthalate, 5, 11 benzonesulphonate, 10 m-nitrobenzenesulphonate, 10 1:4 2-bromonitrobensenesulphonate, camphorsulphonate, 8-

hydrocynaphthalene-1-sulphonate 8 and pyromucate 11
Yttrium silicate.—By heating a mixture of yttria (3 parts), silica (1 part), and calcium chloride (30 parts) for two hours in a wind furnace and extracting the mass with water, Duboin 12 obtained monoclinic crystals of yttrium silicate, Y,O, SiO, which in crystalline form and optical properties closely resembled the mineral gadolinite

Pratt and James, J. Ams. Chem. Soc., 1911, 33, 188
 Cleve and Hogland, Bull. Soc. chem., 1873 [n.], 18, 193, 289.
 Cleve thid., 1874, [n.], 21, 314
 Benner, J. Imer. Chem. Soc., 1911, 33, 50
 Pratt and James, thid., 1911, 33, 1330
 Whittemore and James, thid., 1913, 35, 127.
 Katz and James, thid., 1913, 35, 872
 Errimann and Witth. Annales, 1908, 361, 190.
 Jantsch and Grunkraut, Zedisch. anorg. Chem., 1913, 79, 305
 Holmber, thid., 1907, 52, 83
 Holmber, thid., 1907, 52, 83

<sup>Holmberg, ibid., 1907, 53, 83
Tanata and Voljansky, J. Russ. Phys. Chem. Soc., 1910, 42, 586.
Dubom. Compt. rend., 1888, 107, 99
Grant and James, J./Amer. Chem. Soc., 1915, 37, 2652</sup> 

### ERBIUM 1

Symbol, Fr. Atomic weight, 167 7 (O -16).

Erbium has been found in certain titanium immerals," and is one of the least abundant of the rare earth elements. Erbunn salts are purk or rosered in colour, have a sweet, astringent taste, and exhibit chiracteristic absorption spectra. Certain erbium preparations have been found to exhibit signs of radioactivity 1

The few known compounds of cibium have been described mainly by

Erbium has been described as a dark grey, metallic powder of density 4.77 at 15%. The material was probably impure

Erbium sesquioxide, or erbia, Er.O., is a rose coloured powder of density 8.61 and specific heat thetween 0 and 100 ) 0.065 to 1t does not combine directly with water. A hydrated perocide also exists

Erbium sulphate,  $\text{Er.}(80_1)$ , is described on p. 260, together with the octahydrate,  $\text{Er.}(80_1)$  8H O, which forms monoclinic crystals. The double sulphates, Eng(SO<sub>1</sub>) K<sub>2</sub>SO<sub>1</sub> III<sub>2</sub>O and En (SO<sub>1</sub>) (NII<sub>1</sub>)<sub>2</sub>SO<sub>1</sub> SII<sub>2</sub>O, are readily soluble in water.

Erbium acid selenite, Er O. 18eO, 5H O, is known Erbium tungstate. See p. 265

Erbium silicotungstate.—See p. 266

Erbium nitrate, EreNO ) 5H O, forms large red crystals, stable in an

Erbium platinocyanide, 2(1)(N) 3P((CN) 2HLO is isomorphous with the vitrium salt

Erbium oxalate, Fi  $_3C_2O_4$ ) All O, is a red, increervstilline powder (p. 273). Hydrates with 3, 9, 10, and 14 H O have been described  $^9$ 

Exbrum tormate, 10 mal mate, 11 dimethal phasphate, 1/8 historia naphthal medsulpho acts, 11 and 1/4/2 bromone colorn encsulphomete 15 have also been described

### THULIUM

Symbol, Tin Atomo weight, c 168 5 (O 16)

Thulium is one of the least abundant of the rare earth elements. Puro thulm was isolated by James in 1911, by a method that has been already

- 1 Sometimes known as neo-erbum, since the old "crimin" contained "viterbinin,"

- 1 Sometimes known as neo-erbum, since the old "cribium "contained "viterbunn," holimum, dysposium, and thumina 
  2 K. A. Holmann, Rev., 1910. 43, 2631 
  2 According to Cleve rede infect) the graver part of the "cribium" of Bahr and Bunsen consisted of "ytterbunn" 
  1 Strong, Amer. Chem. J., 1800. 91, 381, et. Cleve and Hogland, Roll. See chem., 1873, [ii...] 18, 193, 289, see also Holmann, los cet. Holmann and Bunger. Rev., 1908, 41, 3783. Holmann and Bunger. Rev., 1908, 41, 3783. Holmann and Bunger. Rev., 1908, 41, 3783. Holmann and Krimicuther, Zeitsch. physikal. Chem., 1910, 71, 312
  2 S. Meyer, Monatch., 1890. 20, 793
  3 Nilson and Petterson. Compet. et al., 1880, 91, 232., Per., 18-0, 13, 1459
  3 Kraus, Zeitsch. Krust. Min., 1901., 34-307
  3 Cleve, lov. ett. (911,0)—Holmann, Rev., 1910, 43, 2631 (344,0) and 1911<sub>2</sub>O), Wirth, 22tsch. annay. Chem., 1912, 76, 174 (1111<sub>2</sub>O)
  30 Cleve, lov. ett.
  31 Erdmann and Wirth, Annalen, 1908, 361-190
  32 Morgan and James, J. Amer. Chem. Soc., 1914-36, 10.
  33 Katz and James, Johl., 1913, 35, 872.

described (p 356). A few compounds of thulium have been described. They are pale green in the solid state and in solution. As the amount of erbium salt present as impurity increases from a trace upwards, the colour of the aqueous solution becomes successively yellowish-green, yellow, colourless, and pink 1

Thulium chloride, TmCl<sub>3</sub>,711,0, is extremely soluble in water and alcohol

Thulium bromate, Tm(BiO), 911,0, crystallises in pale, blush green, hexagonal prisms

Thulium sesquioxide, or thulia, Tm O, has a faint given tuit When carefully made to incandesce, it emits a carmino glow. The oxide dissolves slowly in concentrated acids

Thulium sulphate, Tm<sub>2</sub>(80<sub>4</sub>), and the octahydrate, Tm<sub>2</sub>(80<sub>4</sub>), 811<sub>2</sub>0, have been prepared

Thulium nitrate, Tm(NO<sub>3</sub>), 4H<sub>2</sub>O, is deliquescent and very soluble in water. It may be crystallised from intric acid. In its water content it resembles "ytterbum" intrate

Thulium oxalate, Tm<sub>2</sub>(C<sub>2</sub>O<sub>1</sub>), 6H<sub>2</sub>O, is soluble in aqueous alkali oxalates, with which it forms double oxalates

Thulium acetylacetonate, [(CII, CO), CII] Tai II,O, has been prepared It is not volatile in ranuo

Thulium phenocyacetate,2 cavodylate,3 and 1 1 2 bromonitioben encsulphonate 1 have also been described

### YTTERBIUM

Symbol, Yb. Atomic weight, 173 57 (O=16).

The name ytterbium it now adopted by the International Committee on Atomic Weights to indicate the main constituent of the old "viterbium discovered by Mariginae - Urbain uses the name neoutterbrum, and Auer von Welshach adopts aldebaraneum 9

From the data given by Auer von Welsbach? it appears that 80 to 90 per cent of the old "ytterburn" consists of the new ytterburn. It may be separated from the accompanying element luteenim (or cassiopenim) by fractional crystallisation of the intrates, double ammonium oxalates, or

<sup>James, J. Amer. Chem. Soc., 1910, 32, 517., 1911, 33, 1932.
James, J. Amer. Chem. Soc., 1911, 33, 1932.
Whittemore and James, thid., 1913, 35, 542.
Katz and James, thid., 1913, 35, 872.
This value is due to Urban and Blumentide and was obtained with material free from all but the merest traces of thulums and lateraum. Ancervon Welsbach obtained the value 173 0 m 1913. The atomic weight of lateraum exceeds that of the new ytterburn. To be atomic weight of lateraum exceeds that of the new ytterburn.</sup> therefore a matter of surprise that the determinations of the atomic weight of old "ytterbium," made by Nilson (vide varia) and  $\Lambda$ . Cleve (vide infia), should have given the value 173-1, unless the result obtained by Urbain and Blumenfeld is too high. From the chemical point of view, however, the results of the last named chemists ment the most confidence

6 Auer von Welsbach, Angewer K. Akad Wess Worn, 1905, No. 10, Annalen, 1907,

<sup>351, 464 .</sup> Monatch , 1906, 27, 935 , 1908, 29, 181 , 1909, 30, 695 , Sottongsber K. Akad. Wiss. Wien, 1906, 115, 11 B, 737 , 1907, 116, 11.B, 11.25 , 1909, 118, 11 B, 507 , Urbain, Complement, 1907, 145, 759 , 1908, 146, 106 , Urbain Bourion and Maillaid, dud. ,

<sup>&</sup>lt;sup>7</sup> Auer von Welsbach, Monatsh , 1913, 34, 1713.

bromates,1 the ytterlaum salt being in each case less soluble than the other By the intrate method, Urbani and Blumenfeld? isolated eight successive fractions which furnished specimens of rare earth practically identical with respect to their magnetic susceptibilities and are spectra. According to these experimenters, Auer von Welsbieh's aldelerammin contains some thulium.

Ytterbia, or neogiterbia, Yb,O, is a colomless oxide the mignetic susceptibility of which is  $336 \times 10^{-6}$  e.g.s. units per unit mass. Its salts with colourless acids exhibit no selective absorption. The sulphate terms an octahydrate, Ylc (SO<sub>4</sub>) SH<sub>2</sub>O

The salts of old "ytterlaum" have been examined in detail by Astrid Cleve, but will not be described here !

### LUTECIUM

Symbol, Lu Atomie weight 175 n 1 (O 16)

The name luterium is adopted by the International Committee on Atomic Weights to indicate the constituent of old "vtterbinii that has a higher atomic weight and is less abundant than the new ytterbium The name luterum is due to Urbam, Auer von Welsbach adopts the name cassionerum

Pure luteria has perhaps been isolated by Aner von Welsbach (1913)—It is a colombess oxide, much less priamignetic than ytterbact. Its salts with colombes acids are themselves colombes and exhibit no selective absorption. The anhydrous chloride is more vol (tile than ytterbuum chloride The sulphate torms an ortahydrate, Ln.(80p. 811 0 °

## CELTIUM

Symbol, Ct Atomic weight, (4)

In endeavoiring to isolate lutecium from a large quantity of the immeral galblinite, Urbain obtained a final noncrystalli-alde mother liquor in the course of fractionating the mixed intrates of ytterbinin and Interim. From this liquor he isolated the rare earth present and purified it from all but luteenini and negligible traces of scandium, calcium, and magnesium

The earth was quite white. Its magnetic susceptibility was only onefourth that of nearly pure Interia obtained from venotime, and its arc

<sup>1</sup> See Chapter VI 2 Urtern and Blumenfeld, Compt. rend., 1914, 159, 523 (Trenn and Blumenfeld, Compt. rend., 1914, 159, 523 (Astrid Cleve, Adsch. amorg. Chem. 1902, 32, 129., Chem. New., 1992, 86, 248., 86 also Marignae, Leeh See phus. net., 1878, 64, 57., Compt. rend., 1878, 87, 578. Nilson, 150d., 1879, 88, 642–1880, 91, 56, 118. Fee, 1879–12, 571., 1889, 13–149., Mattynon, Ann. Chem. Phus., 1906, [vm.], 8, 449., Bourron, Ind., 1919, [vm.], 20, 547., 21, 49., Compt. rend., 1907, 145., 243., Kate. and Junes. J. June., Chem. See, 1913., 35, 872., Morgan and James, that., 1914, 36, 10., Janger, Proc. K. Akad. H. etensch. Instrudent, 1914, 76, 1905.

Auer von Welstrich's value (1913) for cassionerum

Aner von Welstich's value '1913) for assequents
 Neoytterba is four to live times as paramagnetic as lutera el'iban, Compt. istol., 1908, 146, 106.
 Mayer, Month., 1908, 29, 1017.
 Aner von Welstich, Anz. gir K. Aleid. Bros. Bros., 1905, No. 10. Aconden, 1907, 251. 164; Month., 1906, 27, 235. 1998, 29, 181. 1669. 30, 695. 1913. 34, 1713; Urban, Compt. rend., 1907, 145, 759. 1908. 146, 106. Urban, Bontron, and Mathad, that, 1909, 149, 127. Urban and Blindienfeld, that., 1914, 159, 323., Bontron, Ana. Chim. Phys., 1900, Ivid. 20, 557. 21. 19. 1910, [viii ], 20, 547, 21, 19

spectrum showed, besides the lines of lutecium, a number of new lines, several of which wore very intense (see p 307).

It thus appears extremely probable that gadolinite contains a new rare earth element that is not present in xenotime. To this element, defined by the are spectrum given on p 307, Urbain has given the name cellium

Anhydrous celtum chloride is more volatile than luteoum chloride, but less so than scandium chloride. Celtum hydroxide is a weaker base than luteoum hydroxide, but is stronger than the hydroxide of scandium. In other respects celtum is also intermediate between luteoum and scandium.

<sup>1</sup> Uthain, Compt. rend , 1911, 152, 141

### CHAPTER XV.

# ACTINIUM AND ITS DISINTEGRATION PRODUCTS.1

Occurrence Actinium is a constant constituent of uranium minerals According to Boltwood, 2m manning muci ils in which radioactive equilibrium has been established the total a-ray ionisation due to the mamnin radium series of products is 1.36 times as great as that due to the uranoun itself, whilst there is an additional activity, equal to 0.28 times that of the manning due to actinium and its products. It follows that actinium is very scarce even in comparison with radium

History.-In 1899, shortly after the discovery of a dimm and polonium in pitchblende, Debicine discovered a third radioactive element in the same mineral. In 1900 he named this element actinium. Deberne found that m working up pitchblende residues, the actimum, which was not precipitated by hydrogen sulphide in acid solution, separated with non and the rare earth elements in the ammonia precipitate, and his final actinium preparations consisted mainly of thorum. For several years no more work on actuating was published by Debierne, and meanwhile Gresel? atmosme d the disenvery in pitchblende of a new radioactive element emonium, characterised by the fact that it gave rise to an emanation which rapidly decayed. Gir el obtained the new element free from thornum, but mixed with the ceruim group of rare earth elements. In 1903 and 1904 Debierne agave further details concerning actinium, described the cinamation to which it gave rise, and declared that emanium and actinium were identical. This identity, at first disputed, was confirmed by Halm and Sackin's in 1905. With the subsequent discovery of the radio-element ionium, it was seen that both Debierne's and Giesel's actinium preparations must have been containenated with that element a

Preparation. The element actionin has not been isolated, meither have actiming salts been obtained in anything approaching a pine state

<sup>&</sup>lt;sup>1</sup> For further information, see Rutherland, Rusermeter systemaces and their Radiations (Cambridge University Press 1912), Soddy, The Chemistry at the Radia are a discharged and the plant of the Annual Reports from 1904 onwards

<sup>2</sup> Boltwood, Amer. J. Sec. 1908, [w.] 25, 269

<sup>3</sup> On the occurrence of a timour products in the atmosphere, see Kurz, Alb. K. Akad.

\*\*Rise Manchen 1909 25, 5

On the occurrence of a finding products in the atmosphere, see Ku17, 15th. K. Akad.

B'iss Manchen, 1909, 25, 5

Deluctive (tough rend, 1809, 129, 593, 1909, 130, 906).

Gresel, Ber., 1902, 35, 3608, 1903, 36, 342, 1904, 37, 16, 6

Debutine, Compt. rend, 1903, 136, 446, 671, 1904, 138, 411, 139, 538

Gresel, Ber., 1904, 37, 5963, 1905, 38, 775; W. Marckwald, Ber., 1905, 38, 2264;

J. Debutine, Physikal Zeitsch, 1906, 7, 14

Hahn and Sackur, Ber., 1905, 38, 1943.

Boltwood, Amer. J. Nat., 1908, [iv.], 25, 365.

Actinium preparations consist for the most part of compounds of the rare earth elements containing unweighable amounts of actumum. Since, moreover, the reactions of actimum are not very well known, strongly active actinium preparations are difficult to prepare and in consequence are extremely expensive. Their preparation from patchblende, carnotite, and the complex Olary ores (S. Australia) may be indicated 1

(i.) In working up prtehblende? for mannum and radioactive preparations, it has usually been the custom to roast the crushed ore with sodium carbonate and then to extract the mass first with witer and then with dilute sulphuric The insoluble residue consists of silica and the sulphates of lead, bismuth, calemin, barmin, rare earth elements, etc. It is heated with a boiling solution of caustic soda, washed and treated with hydrochloric acid, The radium is then found in the insoluble residue and the polonium and actimum in the solution (ii) Numerous processes have been devised for working up the mineral carnotite for manium and vanadium. These have been examined by Plum3 to see which are suitable for extracting also the radio elements it contains. Working with the concentrates from a Colorado carnotite (U -- 4.27, V = 5.5 per cent ) he finally preferred to mix ten parts of carnotite with four parts of anhydrous sodium carbonate and twenty parts of water, heat to boiling for several hours, stirring frequently and adding more water as the mass thickened, and then while hot to filter inder suction, wash the residue with hot water and boil it for eight hours with about five and a halt parts by weight of concentrated hydrochloric acid diluted with ten parts of water. The filtered solution contains the greater part of the radium, polonium, and actinium present in the carnotite (iii) The Olary uranium ore 1 occurs in a lode formation of magnetic titamiferous iron, magnetite, and quartz, in association with biotite. The ore is crushed and then concentrated magnetically. The concentrates amount to 30 per cent of the ore and have the following composition

```
CaO = PbO = 1 e (O_1 - 1) e O = MnO = PbO_3, Ce (O_1) (Ta_0 (D_1, Y)_2 O = Ct (O_3 - U_1O_3 - V_2O_5) - 1)O_3 = S_1O_2 = S_1O_3
   0.55 0.16 17.4 16.9 trace
  3 27
   0.85 | 1.6 | 0.86 | 15.85 | 12.70
```

It is only necessary to decompose about half the ore, by fusion with sodium hydrogen sulphate (salt cake), to seeme a practically complete yield of the radioactive contents. The fused product is crushed, agriated with water, and the finely divided silica and lead, barrum and radium sulphates separated from the coarser, unchanged ore by clutration. The turbid liquid is allowed to stand and the "slime" of crude sulphates and silica separated from the clear liquid, which is treated for mamum. The manum liquor contains part

<sup>&</sup>lt;sup>1</sup> In consequence of the tayless nature of the initial change of actinium, it often happens that actinium preparations when separated from innerals are scarcely radioactive, then activity and emanating power, however, increase enormously during the next few months

activity and chanating power, however, increase enomionsly during the next few months. It is thus easy to overlook the presence of actinium, hence, in seeking to obtain actinium preparations no material should ever be thrown away in a hinty.

2 Dehierne, Compt. rend., 1899, 129, 593, 1900, 130-906. Mine Curie, Theses (Pairs, 1903), or translation in them. News, 1903, 88, Haitinge and Ulrich, Nit winder A. Range Viss. Wen, 1908, 117, in a, 619, Monalsh, 1908, 29, 485, Anei von Welskerh, St. wags ber. K. Akad. Wiss. Wich, 1910, 119, in a, 1, Monalsh, 1910, 31, 1159, Zeitsch. award, Chem, 1911, 69, 353, J. So. Chem. Ind., 1911, 30, 535, Boltwood, Proc. Roy. Soc., 1911, A, 85, 77, Le Radium, 1911, 8, 104

3 Phun, J. Amer. Chem. Soc., 1915, 37, 1797

4 Radeliff, J. Roy. Soc. Acta South Bules, 1913, 47, 145, 1914, 48, 408, Mining Sec., 1914, 69, 37, J. Soc. Chem. Ind., 1914, 33, 229, Chem. News, 1915, 111, 59

of the rare earths originally present in the ore, but practically none of the actinium, which is found, along with the remainder of the rine earths, in the "shme

It will be seen that pitchblende residues and Olary ore "slimes" are somewhat smalar in composition. Radchil originally worked up the latter in the following manner. The 'slimes" are treated with sulphuric acid, filtered, and boiled with excess or an aqueous solution of solution cubonate Insoluble sulphates are converted into carbonates and much silica is dissolved The washed earbonates are dissolved in Technochloric acid and the sulplantes again precipitated. The enide sulphates thus obtained are fised with sodium carbonate in graphite pots and the product digested with hot water. The insoluble residue, after picking out the metallic lead, is dissolved in livdro chloric acid and the solution evaporited to dryness to dehydrate the silica The chlorides are dissolved in dilute hydrochloric acid, filtered to remove silica, and the solution is their siturated with hydrogen chloride. Brimm, and radium chlorales are quantitatively precipitated, nearly tree from other elements, whilst the actimum remains in solution with the rare cartles. This method has subsequently been modified as follows. The crude naxture of sulphates is fused in an iron crucible with excess of caustic sodic and a lettle so from carbonate. The melt is repeatedly extracted with hot water, whereby most of the lead is dissolved, and the insoluble residue is digested with sodium carbonate solution under a pressive of 90 Hs, per square inch. The carbonate residue is washed converted intochlorides, freed from silica, and the barroin and radinar separated as before !

Properties - Little is known of the chemical properties of actimum It has been known since its discovery that actinium resembles the rare earth elements. Debicing was in circi, however, in regarding it as inalogous to thorium. It is most similar in properties to the tervalent rare earth When these clement. or the ecrnin group, particularly lanthaming actimierous rare earth elements are fractionated as the double magnesium intrates the actimina concentrates in the neodynamic and samarium fractions 1

Actimium is not precipitated by hydrogen sulphide in dilute acid solution (separation from polonium and radioload), but is precipitated as hydroxide by ammonia (separation from radium). This precipitation, however, is very integrant, and in the presence of considerable quantities of anamonium salts is livenoments complete. Plum, who studied the precipitation of actinium by annuoma in the presence of a little almomonic silt, found it impossible to separate all the actimum from raditerous and actimiterous barium, chloride, in

Ploo, Ruc Larths Bison 1.0 5105 7.05 3 0 to 1 cent  $_{6}q.2$ 3 0 12.52.2 10.8

Ł

 $<sup>^4</sup>$  According to Rad hildo-  $e^+e^-$ ten tous of Olay one concain ites valid only forty kilo grams of crude sulphrites of the foll ving composition -

Debetne, Cos pt. rent., 1899-129 (93), 1960, 130 (96)
 See Greed, Ber., 1967, 40 (601), Strombolm and Svelberg Zeech, anoral Chem. 1909, 63, 197

<sup>1909, 63, 197

\*</sup> Debette, Complete ad., 1904, 139, 5.8

\* Leven, Plat Mag., 1906, [ver], 12 177, Physikal Zerm., 190, 8, 129, 11 din, Plat Mag., 1906, [ver], 1907, [ver], 13, 165, Boltwood Amer. J. Ser., 1908, [ver], 25, 292, Proc. Roy. Soc., 1911, A, 85, 77, And von Wessleid locate. For remove ammonium salts, evaporate to diviness and heat the residue with boiling intreased or aqualogia, then remove excess of redshe evaporation.

\*\*Plum, J. Amer. Chem. Soc., 1915, 37, 1797

Actinium is not precipitated from its chloride solution by boiling with sodium thiosulphate, nor from its nitrate solution by the addition of hydrogen perovide (separation from thorium and ionium). It is precipitated as oxalate by oxalic acid or ammonium oxalate, in this precipitation the concentration of free mineral and should be reduced to a minimum, since actimum oxalate, like lanthamim oxalate, is decidedly soluble in immeral acids. The fluoride of actinium, like the fluorides of the rare earth elements, is insoluble in water and aqueous hydrofluorie acid (separation from zincomum and titanium) When a little barrum salt is added to an actimum solution and the barmin precipitated as sulphate, the precipitate adsorbs the actimum This property, observed by Debierne, is utilised in separating actinium from pitchblende and Olary ore. Actimum is readily separated with manganese when that element is precipitated from basic solutions as a mangainte 1 Auer von Welsbach describes actumum as being intermediate in chemical properties between lanthamum and calcium

Actimium is an unstable element, and like thorium and uranium, it is constantly breaking down at a slow but definite rate, and so giving rise to a series of radioactive disintegration products. When in equilibrium with these products,  $a_{\gamma}$ ,  $\beta$ , and  $\gamma_{\gamma}$  are emitted by actinium preparations. The a rays are very powerful, preparations 100,000 times as active as manning oxide have been obtained. The  $\beta$ -rays have a relatively small penetrating power, and the y-rays are feeble both in activity and in penetrating power. Tho most striking feature of an actimum preparation is its emanating power (p. 419). In the course of the radioactive transformations occurring in actimini preparations behum is evolved, and aqueous solutions of the preparations slowly evolve hydrogen and oxygen 2

## The Disintegration Products of Activium.

History. —It was early known that actimum produced a rapidly decaying emanation, which gave rise to radioactive deposits upon surrounding objects. The modern theory of the atomic disintegration of radioactive elements was put forward in 1903. In terms of this theory, Rutherford, in the Bakerian Lecture for 1904, gave the actinium series as follows

Element . . 
$$\Lambda e \Rightarrow \Lambda e \chi \xrightarrow{\alpha} \frac{\alpha}{\beta}$$
  $\beta$  Element . .  $\Lambda e \Rightarrow \Lambda e \chi \xrightarrow{\beta} \Lambda e Em \Rightarrow \Lambda e \chi \Rightarrow \Lambda e B \Rightarrow \Lambda e C$  and  $3.7 \text{ Secs.} 41 \text{ mms.} 1.5 \text{ mms}$ 

The existence of actinium-X was assumed by analogy with the nramum and thorum series. The assumption that the emanation gave rise to an active deposit, actinium-A, which was transformed into another element, actinium-B, and that this in turn was transformed into a non-radioactive and stable element, actinium-C, was based upon Miss Brooks' study of the actinium active deposit 1

With the publication of the disintegration theory, the view arose that the α-particles contted in radioactive changes are charged atoms of helium. Although this view was not definitely established until 1908, its probability

Aner von Welsbieh, loc cit

Debierne, Compt. rend., 1905, 141, 383, Ann. Chim., 1911, [ix.], 2, 27, 428.

Rutherford, Phil. Trans., 1905, A, 204, 109

Miss Brooks, Phil. Mag., 1904, [vi.], 8, 173

was quickly recognised when in 1903 Ramsay and Soddy demonstrated the continuous production of beliam from radium. The growth of beliam from actinium was observed by Debicine 1 in 1905 and confirmed by Giesel 2 in 1907

In 1905 Godlewsla<sup>4</sup> and Gresel<sup>4</sup> independently discovered the missing element actinium X (half period = 10.2 days). Godlewski's results indicated that the change from actimum to actimum V is a rayless one. In the following your, however, Hahn? discovered that the rayless change is from actinium to a new element radioactinium (half-period 195 days), which expels a-rays and passes into actinium X

In 1908 Hahn and Meitner? discovered that the transformations of the active deposit produced by the decay of actinium enaurition are more complex than had been intheir supposed. Then views on the degradation of the deposit may be thus summarsed

Element . 
$$AcA \rightarrow AcB \rightarrow AcC \rightarrow AcD$$
 condition that period 36 turns 2.15 turns 5.1 turns

The active deposit, however, has proved to be still more complex, for in the same year Bronson? discovered that the emanation produces is us decay twice as many a particles as does the active deposit it produces. In accordance with the preceding scheme for the degradation of actinium \( \), this means that each atom of emanation evolves two a particles in its decomposition. This fact was confirmed by Geiger and Marsden, in 1910. The explanation was supplied by Gerger" in 1911. The emanation breaks down with the loss of one a particle per atom decomposed into a new solid element which decays with extraordinary rapidity, also losing one a particle per atom decomposing, and becoming transformed into the element previously known as actuming V. Geiger determined the half-period of the new element to be only 0.002 second, and Moseley and Fajans, in who developed a highly accurate method for measuring such a short period, obtained exactly the same result. The new element was called actinium A, the elements previously known as actinium-A, B, C, and D respectively being renamed actinium B, C, D, and E.

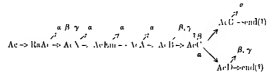
The foregoing, however, does not completely describe the transformations of the active deposit. In 1909 MHe Blanquics 11 concluded that actinium-C (then called AcB) gave rise to two kinds of a rays. Her experimental results have not been confirmed,1 but the experiments of Marsden and others leave httle doubt that actinium C is complex. The a rays it emits have been found 

<sup>1</sup> Debreine, Compt. road., 1905, 141, 383. For full experimental details, we Debreine, Ann. Phas., 1911, [ix.], 2, 4.28.
2 Gresel, Ber., 1907, 40, 3011.
3 Godlewski, Phil. May., 1905. [vi.], 10, 35, 375. Bull. Irad. Ser. Cracow, 1905, p. 265.
4 Gresel, Julius Radionkliv, Lichtmark, 1904, [vi.], 13, 58.
5 O. Hahn, Ber., 1906, 39, 1005., Parl. Man., 1906, [vi.], 12, 244., 1907, [vi.], 13, 165., f. Levin, Mack., 1906, [vi.], 12, 177.
6 Hahn and Mertner, Physikal Zerisch., 1908, 9, 649, 697., Hahn, thad., 1907, 10, 81.
7 Breisson, Phil. Mag., 1908, [vi.], 16, 291.
6 Geiger and Marsden, Physikal Zerisch., 1910, 11, 7.
9 Geogre Phil. May., 1911, 11, 12, 201.

<sup>&</sup>lt;sup>8</sup> Geiger and Matsden, Ingeneral Learner, 1917, 22, 23
<sup>10</sup> Geiger, Phil May, 1911, [1] 22, 201
<sup>10</sup> Mossley and Fajans, that, 1911, [1], 22 529
<sup>11</sup> Mile, Blanquies, Compt. rend., 1909, 148, 1753, 1910, 151, 57. Le Read am, 1909, 6, 230, 1910, 7, 159
<sup>12</sup> Sec, e.g., Moseley and Fajans, loc cit., Vaider and Matsden, vide infra.

to be of two kinds, their ranges in air being 5.4 and 6:15 cms., the latter only arise from 0.15 to 0.20 per cent of the atoms undergoing change. The explanation is as follows - Actimum C breaks down in two different ways, in one of which an a particle of range 5.4 cms is evolved and actinium-D produced, while in the other and slower method a  $\beta$ -particle is evolved and a new element, actinium C', is produced The relative amounts of AcD and AcC' produced are as 99 83 is to 0.17, and the new element decays at almost exactly the same rate as actimini-A, with the emission of a-rays of range 6.45 cms

In 1912 Chadwick and Russell<sup>2</sup> announced the existence of an element intermediate between radioactinium and actinium-V. This was denied by Hahn and Meitner,3 and subsequently Chadwick and Russell withdrew their claim. The actimum series, as it is at present known, is accordingly as indicated in the following scheme, in which the nature of the radiation from each product is also shown -



The  $\alpha$ ,  $\beta$ , and  $\gamma$ -radiations. Actimum, when carefully freed from its products, is found to emit no detectable  $\beta$ - and  $\gamma$ -rays, and to emit  $\alpha$ -rays to so slight an extent that the phenomenon is presumably due to traces of its products. In other words, the change from actimum to radio atmium is a rayless one Six actimini products, viz radioactimini, actinium-A, actimini cinamation, actimini-A, actimini-C, and actimini C, decay with the emission of a rays, one a-particle being expelled per atom decomposed The ranges (in cms) of these a rays in an at NTP are given as follows by varions experimenters - "

|                         |              | -                |      |      | · -  |      |
|-------------------------|--------------|------------------|------|------|------|------|
|                         | Rive         | AcX              | Aclm | 101  | AcC. | Acc' |
|                         |              |                  | _    |      |      | ;;   |
| Geiger and Nuttill?     | 4 :50        | 1 17             | 5 10 | 0.16 | + 12 | ļ    |
| Meyer, Hess, and Paneth | 1 0 and 1 37 | 4 (1)            | 5 25 | 5.94 | 1.88 | 1    |
| Vaider and Marsden"     | ,            |                  |      |      | 51'  | 6 11 |
| M'Coy and Leman " .     | 1 17         |                  |      |      |      | i    |
|                         |              | F. Communication |      |      |      | 1    |
|                         | -            |                  |      | -    |      |      |

Marsden and Wilsen, Nature, 1913, 92, 29.
 Marsden and Perkins, Phil Mag., 1914, [vr], 27, 690.
 Vaider and Marsden, ind., 1914, [vr], 28, 818.
 Chadwink and Russell, Nature, 1912, 90, 463.
 Hahn and Mettner, Phinshal Zeitsch., 1913, 14, 752.
 Hassell and Chadwick, Phil Mag., 1914, [vr], 27, 112.
 Hahn, Ber., 1906, 39, 1605.
 Phil Mag., 1907, [vr], 13, 165.
 Lee, in that, 1906, [vr], 12, 177.
 Hahn and Rothenbach, Physikal Zeitsch., 1913, 14, 109.
 According to S. Meyer, Res., and Paneth (vide infia), however, actinium cinits a rays of range 3-38 cms in 4n at N.T.P.
 The range in air is directly proportional to the absolute temperature and inversely proportional to the messure.

proportional to the pressure 'George and Nuttall, Phil. Mag., 1912, [v1], 24, 647

8 S. Meyer, Hess, and Paneth, Sitzuanjsbr. K. Mad. Wiss. Wien, 1914, 123, n. a, 1459.

9 Varder and Marsden, Phil. Mag., 1914, [v1], 28, 818

10 M'Coy and Leman, Phys. Review, 1914 [n], 4, 409.

Three members of the series, viz radioactimum, actimum B, and actinium-D, emit  $\beta$ -rays. The nature of these rays has been studied by various physicists. 1 those emitted by radioactiming and actiming B have very httle penetrating power. Associated with the Brays are y-rays. Radio actinium and actinium Demit very penetrating y rays, and a timum Blikewise emits fairly powerful years. Radionetinium and actimum-Balso contrautly weak yrays, while in addition to this actinium B emits very feelib, raivs?

It will be abserved that radioactiming courts both a- and  $\beta$  rays. Only one other radio element is known to do this, viz. radium

Rates of Decay. The various members of the actinium series differ greatly in stability, the half periods varying from years with actinium itself to 18 88 days in the case of radioactinium, and to 0 002 second in the case of actimum A. The precise rate of change of actimum itself is unknown According to Soldly, however, its period of average life probably does not exceed a hundred veris, and Mior Curie, has made observations which point to a period of only thirty years. It should therefore be possible to observe the growth of actimina. The fact that this has never been done suggests the existence of an intermediate long-fixed product between actinium and radioactimium. The matter, however, is still obseine

The radioactive constant (A), half period (T), and period of according life (P) of each of the members of the actimum series are given in the following table (P 1/A 1 11 (T) -

| Fh nænt      | \(\-(\-\) ( | 1          | P          | 1 lenocut   | ' \(\langle (see ) \) | 1          | P         |
|--------------|-------------|------------|------------|-------------|-----------------------|------------|-----------|
|              |             |            | 1          | -           |                       |            |           |
| Rabor (maao) | 4 '5 10 7   | 15 85 days | " "Hidos   | Actonnia B  | 3'1 1'                | nomm !     |           |
| Actinom X5   |             |            |            | Actinama C# |                       | 24 nor 8 1 |           |
| Emonation 6  | 177 16 5    | 3) 11541   | 16 680 8   | Actuana Da  | 2 45 10               | 1.160      | 0 9)11(08 |
| Actioned 🐷   | 16 ق        | 0.002.40   | 1 1813 644 | l           |                       | :          |           |

In the manimi and thorium series the following empirical relationship has been discovered, the longer the period of the element the shorter the

See H due and Mether: P<sup>2</sup>ys, lat Zet k. 1208, 9. n [9-607] O conclusiver, Halin, and Mether, x, t. 1913, 14 (52), yr. Goldwski, P. & Maa, 1905, [vi l. 10-35, 475].
 Levin, x, d., 1906, [vi ], 12, 177
 See Ratherlord and Richarlon, Phot. Max. 191 [vi ], 26, [6] d. d. f. and W. M. Saddy and Rickli, bold, 1910, [vi ], 19, (25). Rocall and F. Soddy, that, 1911, [vi ], 21, 130. Russell and Chadwock, that. 1911 [vi ], 27, 112
 Mon. Curo. Ir. Reviews, 1911-8, 3
 M. Coy and Lamin, Irans. R. See. 1914, [vi ], 4, 409. Sef. Halin, Ib. I. Maa, 1907, [vi ], 33-165. Helm and Robbindock, Prop. Int. Z. Sec. h. (2011) 14, 160
 M. Coy and Laman, Physical Zet Sec. 1913, 14, 1280, and sec. t. ci. Goddewski, Phot. May. 1905, [vi ], 10, 35. Gersel, Rev., 1907, 40, 3011. Helm and Robbindock, Proc. etc.

be eet.
 b P B Perkins Pred Mag , 1914, [vi ], 27 720 , et 15 Jaccine, tomplere al., 1904 138, 411 , Halme and Suckar, Bre , 1995, 38 (1915). Mr. M. Leslie, Phei Mag. 1912, [vi ].

<sup>24, 637</sup> <sup>7</sup> Moschy and Fajans, Phil. May. 1911, [ve] 22, C29, Geiger, dod, 1911, [ve],

<sup>22, 201</sup> 8 Halac and Mechaer, Physikal Met. 4, 1908, 9, 649 of Mr. Broxl. Phil Man., , 1904, [vr.], 8 373 Hess Sit warder, K. Mont. Hast. Han. 1904, 116 n. a., 1121, 8 Meyer and von Schwendler, Pold, 1905, 114, n. a. 1147 Bronson, Amer. J. Sc., 1905, [vv.], 19, 185 "Hahn and Mertner, live cit., ct. Miss Brooks, live et.

<sup>19</sup> Kovářík, Physikal Zedsch , 1911, 12, 83 , c/ Hahn and Meitner, loc est

range of its  $\alpha$ -particles at NT.P, and if the logarithms of the ranges are plotted against the logarithms of the radioactive constants or periods the points he on straight lines in each series (the Geiger-Nuttall Law). To this rule in the actinium series radioactinium constitutes an exception if Geiger and Nuttall's values for the ranges are accurate, according to M'Coy and Leman, however, their value for radioactinium is too high, while Meyer, Hess, and Paneth state that this element gives  $\alpha$ -rays of two different ranges, an indication that the actinium series is more complicated than it is at present thought to be Applied to actinium C', the rule indicates that the values of  $\lambda$ , T, and P must be almost identical with those for actinium-A

From the periods here given it is possible to calculate the manner in which the  $\alpha$ - and  $\beta$ -ray activities of actimum, radioactimum, and actimum X, initially free from any other radioactive elements, vary with the time, provided the radioactive preparations lose none of their radioactive products through loss of gaseous emanation, experimental and calculated results are m good agreement. The a-ray activity of actimum, initially zero, develops as the preparation is kept, slowly at first, then more rapidly, and finally more slowly again, reaching a limiting value after four months, when the actinium and its products are in radioactive equilibrium 1. With radioactinium, the a-ray activity uses to a maximum of about 2.25 times the initial value in seventeen days and then dimmishes, the rate of decay soon approximating to the exponential value for radioactinium and being practically complete in four months The  $\beta$ -ray activity varies similarly, its maximum being reached in about twenty days,  $^{1,2,5}$ . The  $a_{\gamma}$  and  $\beta$ -activities of pure actinium- $\lambda$  rise to their maximum values in about four homs, and then decay at a rate which ultimately becomes exponential with the period of actinum-X.2

Separation of the Members of the Actinium Series. - Radioactinium may be separated from its parent and its products by adding a little thorium salt to an acid solution of the actinium preparation and borking with an excess of sodium throsulphate. The precipitated thorium basic throsulphate carries with it the radioactinium. Hahn, who originated this throsulphate method, added no thorium salt, and he described the separation as uncertain, probably his actinium contained a little thorium. To ensure the purity of the radioactinium, the precipitate may be dissolved in hydrochloric acid and reprecipitated by sodium throsulphate. Instead of adopting the throsulphate method for separating the thorium and radioactinium, the hydrogen peroxide method (p. 320) may be used 68. If the addition of thorium is undesirable, a little zirconium may be added and the throsulphate method employed. By fractionally precipitating an actinium solution with ammonia, radioactinium concentrates in the first fractions.

Actinium-X may be separated from a solution of an actinium preparation by precipitation with ammonia, which leaves the actinium-X in the filtrate 19

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Hahn, Phil Man, 1907, [v1], 13, 165
Hahn and Rothenbach, Physical Zertsch., 1913, 14, 409
M'Coy and Leman, that, 1913, 14, 1280.
Levin, Phil Mag, 1906, [v1], 12, 177.
M'Coy and Leman, Phys. Renew, 1914, [u1], 4, 409.
Fleck, Trans. Chem. Soc., 1913, 103, 381.
Hahn, Ber., 1906, 39, 1605; Physikal Zertsch., 1906, 7, 855, Phil. May, 1907, [v1], 13, 165.
M'Coy and Leman, Physikal. Zertsch., 1913, 14, 1280
Hahn and Rothenbich, that, 1913, 14, 409
Godlewski, Phil. Mag., 1905, [vi.], 10, 35.
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To ensure the removal of actimum and radioactimum the filtrate may be acidified and reprecipitated by ammonia after the addition of a little ferric The filtrate is then evaporated to dryness and the residue ignited to expel the actiminm "active deposit," or the actimina-X is removed from solution by co-precipitation with barium sulphate? Freedom from actinium is probably best ensured by growing the actimum A from pure radioactimum, and after a suitable interval of time removing the residual radioactinium as has been previously described 3

Actinium emanation may be removed from actinium solutions by passing a rapid stream of an or other gas through it. The emanation is also evolved with great facility from most solid actimum preparations, particularly the The remarkably rapid rate of decay of the emanation and the readiness with which it is evolved make it easy to demonstrate the emanating power of an actinium preparation by merely holding it over a large zinc sulphide screen in the dark. The emanation streams away and illiminates the screen, and as the emirents in the air blow the emanation about so the screen becomes illuminated in different places

Actinium active deposit — In view of the excessively brief life of actinium  $\Lambda_i$ it cannot be isolated and studied like an ordinary radio element. In all ordinary work, actiminic emanation and actiminia V act together as one product. By then decay, actimini B, C, C, D, and the end products arise Collectively they are termed the "active deposit," since they are produced on all objects with which the cinanation comes into contact. To separate tho active deposit a current of an laden with emanation may be passed through a metal cylinder provided with a metal electrode held axially in position and insulated from the cylinder by rubber stoppers. A potential difference of about 50 volts is maintained between the central electrode and the cylinder, the latter being positive to the former. The active deposit then collects on the negative electrode. The yield is neve-more than 95 per cent 4

The manner in which the a ray activity of the "active deposit" varies with the time depends upon how long is taken to collect the deposit. If this does not exceed ten seconds, the activity increases slightly with the time, reaches a maximum in seven munites, and then decreases, the decay soon becoming exponential with the period of actinium-B, the longest lived element in the deposit. If the time taken to collect the deposit exceeds ten seconds the initial use in its o ray activity cannot be observed?

Actimum B is more volatile than actimum C. When the active deposit is collected on a platimum wire and heated in air, actimum B begins to volatilise at 400° C, and is completely volatilised in ten minutes at 750°C, b in four minutes at 900' C, or in about half a minute in the blowpipe flame, when actimism-D is also volitilised. Actimism C is not perceptibly volatile below 700° C 6 After having been volatilised in air, actiminin-B becomes

<sup>1</sup> Hahn and Rothenbach, Physikal Zeilsch., 1913, 14, 109

J. Hahn and Rothenbach, Physikal Zedych, 1913, 14, 109

M. Coy and Leman, Phys. Review, 1914, [n.], 4, 109

M. Coy and Leman, Physikal Zedsch, 1913, 14, 1280

M. Coy and Leman, Physikal Zedsch, 1913, 14, 1280

M. Coy and Leman, Physikal Zedsch, 1913, 14, 1280

M. On the behaviour of the active depens in an electric field, see Miss Brooks, Phil Mag, 1904, [v.], 18, 373, Russ, thid, 1908, [v.], 15, 601, 737, Kennedy, thid, 1909, [v.], 18, 744, M. Leman, thid, 1912, [v.], 24, 370, Walnebey thid, 1913, [v.], 26, 381, Lamax, thid, 1914, [v.], 28, 761, Amer J. See, 1914, [v.], 38, 539.

Miss Brooks, Phil. Mag, 1901, [v.], 8, 373

S. Meyer and von Schwedler, Sitzungsber K. Akad Wiss Wien, 1905, 114, in. a, 1147, Levin, Physikal Zedsch., 1906, 7, 812, Hahn and Meitner, thid, 1908, 9, 649; Schrader, Phil Mag, 1912, [v.], 24, 125.

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much less volatile and may be condensed on a surface at 1000° C.; this is presumably due to oxidation of the element, since the phenomenon is not observed in an atmosphere of hydrogen. Laposure of the active deposit for ten to fifteen minutes to bromine vapour increases the volatility of actinium-B, as also does exposure to chlorine or hydrogen rodide. The last two gases, however, cause actinium-C to become more volatile; in fact, after treatment with hydrogen iodide, actinium-U is more volatile than actinium-B. Exposure to hydrogen chloride does not affect the volatilities, but if the active deposit be then treated with water, nearly all the actinium-B is dissolved, but no actimum-C Actinium-B is very readily dissolved by dilute acids; immersion of the active deposit for one minute in 0 001N hydrochloric or sulphuric acid causes 80 to 95 per cent, of the actimum-B to be dissolved

and practically none of the actimum-C <sup>1</sup>

The actimum active deposit may be dissolved in boiling hydrochloric acid. By electrolysing this solution between platmum electrodes, part of the actinium-C may be deposited on the cathode. A much better and simpler method of separating the actinium-C from this solution is to have it nearly neutral, heat it to boiling and immerse a piece of sheet nickel in it for a few imputes. The actimum C is deposited on the nickel Actimum-D may be withdrawn from the solution by shaking with animal charcoal, in which it is adsorbed.4 It is best prepared, however, by the method of radioactive recoil.

An atom of actimum-C, of mass 210, loses an a-particle, i.e. a charged atom of helium, of mass 4, when it disintegrates. The helium atom is expelled with a velocity of  $1.89 \times 10^9$  cms. per second, and in accordance with the second law of motion the residual atom of actimum-D, of mass 206, must recoil (just like a gun recoils when it discharges a projectile) with a velocity of  $(4\times1.89-10^9)/206$  or  $3.67\times10^7$  cms. per second. The actinum-1) as it is produced has accordingly a kind of "nuscent" volatility and feeble penetrating Since the a-particles are discharged in all possible directions, so also do the atoms of actinium-D recoil in all directions, and if an excessively thin layer of active deposit is collected on a smooth, carefully polished metal surface, 50 per cent of the actimum-D atoms produced in its decay may be expected to recoil normal to and away from the plate. The recoiling particles carry positive charges and deposit preferentially on a negatively charged surface. Accordingly, by placing the metal plate carrying the active deposit opposite the surface on which the actinium-I) is to be collected, both being contained in an evacuated vessel, and establishing a suitable potential difference between active deposit and receiving surface, it is possible under good conditions to collect almost the theoretical yield of actinum-D 5

Physical Properties of Actinium Emanation.—Actinium emanation is a gas which so far as is known does not enter into any chemical combination. At -120° C. it begins to condense, and condensation is

¹ Schnader, loc ct.
² Miss Brooks, Phil. Mag., 1904, [vi.], 8, 373.
³ Meither, Physikal. Zeitsch., 1911, 12, 1094.
⁴ Hahn and Meither, that, 1908, 9, 649.
⁴ Hahn and Mether, loc. ct.; Kovářík, Phil. Mag., 1912, [vi.], 24, 722, Wood, ibid, 1913, [vi.], 26, 588; Flack, Trans. Cicim. Soc., 1913, 103, 1052. The recoil method is a general one for preparing the products of a-ray changes. Some curious early observations on the decay of actinum active deposit (S. Meyer and von Schwedler, Stitungsber, K. Akad. Wiss. Wien, 1905, 114 · ii a, 1147, 1907, 116, ii, a, 315) were ultimately traced to contamination with actinum-X which had recoiled at the moment of its formation (Hahn Physikal. Zeitsch. 1909, 10, 81). Physikal. Zeitsch., 1909, 10, 81).

complete at -150° C, agreeing in these respects extremely closely with thorium emanation. It is twice as soluble in water as thorium emanation, and six times as soluble as radium emanation. The following, as solvents for actinium emanation, are arranged in increasing order of absorbing power, saturated potassium chloride solution, water, sulphuric acid, alcohol, ammonia, acetone, benzaldehyde, benzene, toluene, petrolemin, and carbon disulphide. The emulation is strongly adsorbed by cocounit charcoal at the ordinary temperature

The rate of diffusion of the chamation in various gases has been determined by several observers with results of doubtful value from the point of view of calculating the molecular weight or the emanation? It is very improbable that the ordinary laws of gaseous diffusion can be applied to the case of an excessively minute amount of characteristic diffusing into a relatively enormous amount of another gas. From comparative experiments on the actinium and thorum emanations (in the carrying out of which various sources of possible error in the work of previous experimenters were detected), Miss Lashe | concluded that these two emanations have molecular weights that are nearly equal, and it is known that the value for thorium emanation is 220.2 . Marsden and Wood's have determined the nodecular weight of actinium emanation by a new diffusion method in which gas other than the emanation is present only in minute amounts. Then experiments lead to a value of about 230 for the molecular weight, and this result is probably tairly accurate, since Debicine has devised a method which is based upon the same theoretical principles as that of Marsden and Wood, and by its use obtained a value of 221 for the molecular weight of radium emanation, the correct value being otherwise known to be 222

Chemical Properties of the Members of the Actinium Series. --The account already given of the preparation and properties of actir um leaves little doubt as to the tervalency of that element. In 1909 Stromholm and Svedberg? examined the chemical nature of various radio elements by crystallising salts from their solutions and seeing whether the radio elements separated with the crystag or not - In this manner they found that actinium was isomorphous with lanthammi, actimum V with the alkaline outly metals, and radioactmum with thorum. A physico-chemical method for determining the valencies of the radio elements was devised in 1913 by you Hevesy 8 who found that actimum is tervalent and actimum-X is bivalent

The chemical properties of actimum and several of its products have been carefully examined by Fleck," with the following results. Actinium is so similar to mesothorium 2 that when once mixed, chemical means fail to effect any separation of one from the other Similarly, radio actimina is chemically in distinguishable from thorum, and actimum B, actimum C, and actimum D are identical in chemical properties with lead, bismuth, and thallium respectively.

The chemical identity of two elements having different atomic weights is

Kinoshita, Phil. May, 1998, [vi], 16, 121. of Henriot, Le Radium, 1908, 5, 41.
 Von Hevesy Physikal Leits h. 1911, 12, 1214
 Debierne, Le Radium, 1907, 4, 213. Bruhat, ibid., 1909, 6, 67, Compd. rend., 1909, 148, 628. Russ, Phil. May, 1909, [vi] 17, 412
 Miss Leshe, Phil. May, 1912, [vi] 24, 637
 Massden and Wood, Phil. May, 1913, [vi], 26, 918.
 Debierne, Low. Phys. 1915, [vi], 26, 918.

<sup>Matsden and Wood, Phil. Maj., 1915, [vi], 20, 916.
Debierne, Ann Phys., 1916, [ix ] 3, 62
Stroibloin and Sveilberg, Zeits h among Chem., 1909, 61, 338; 63, 197.
Von Hevesy, Phil. Mag., 1913, [vi.], 25, 390, 1914, [vi.], 27, 556.
Fleck, Trans. Chem. Soc., 1913, 103, 381 (Ac, RaAc, AcB, AcC), 1052 (AcD).</sup> 

a very startling discovery. The preceding results are only a few of those that have been experimentally investigated. A group of elements which are chemically identical is called a group of isotopic olements. The isotopic groups, so far as they have been experimentally examined and relate to members of the actinium series, are as follows —1

- (1) Radioactimum, thorum, radiothorum, ionimi, and mannim-X.
- (n ) Actinium and mesothorium-2
- (m) Actmum-X, tadmm, mesothorum-1, and thorum-X.
- (iv ) Actimum emanation and the emanations of radium and thornin.
- (v) Actinium-B, lead, radium-B, thorum-B, and radium D
- (vi.) Actinum-C, bismuth, i.dium-C, thorum-C, and radium-E.
- (vii.) Actimum-D, thallium, and thorium D

The identity of isotopes extends to their electro-chemical behaviour. Actinium-B and actimium C, for instance, are electro-chemically identical with the B and C members of the thornum and radium series?

With the recognition of isotopism it became possible to assign the radioelements positions in the Periodic Table, for it was seen that two general rules could be framed (i) An element formed in an a-ray change (i.e. a mass change) differs in valency from its parent by two, and so occupies a position two places behind its parent (i.e in the direction of diminishing mass). (ii) An element formed in a  $\beta$ -ray change (i.e. no change in mass) differs in valency from its parent by one, the valency increasing in the change, the product accordingly occupies a position one in advance of its parent these rules the members of the mamum and thorum series may be placed in position since the starting-points for thorium and uranium are known, and it is found that a group of isotopes occupies one space in the Periodic Table In the absence of definite knowledge concerning the origin and atomic weight of actimum, the actimum series must be placed in position in accordance with the principle of isotopy, and it must further be assumed that the rayless change  $Ac \rightarrow RaC$  follows the  $\beta$ -ray rule <sup>3</sup>. The actinium series then falls into Mendeléeff's table as follows -

|                                 |            | -  |           |           |            |           |             | , |
|---------------------------------|------------|----|-----------|-----------|------------|-----------|-------------|---|
| -><br>Group                     | 0 ,        | 1  | 2         | 3         | 4          | 5         | б           | 7 |
| Fourth long period (Odd series) | - i        | Λu | Hg        | Tl<br>AcD | l'h<br>AcE | Bı<br>AcC | AcA<br>AcC' |   |
| Fifth long period (Even series) | Nt<br>AcEm |    | Ra<br>AcX | Ac        | Th<br>RaAc |           | U           |   |

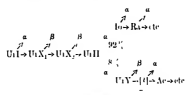
Origin of Actinium.—Actinium is a constant constituent of uranium minerals, and it occurs in them in amounts proportional to their uranium contents 5 It is therefore hard to resist the conclusion that actinium is

Soddy, The Chemistry of the Radio elements (Longmans & Co., 1914) pt. 11, p. 14.
 Von Hevey, Phil. Mag., 1912, [v1], 23, 528.
 This has also to be assumed for the rapless change MsThI→MsThII.
 A short account of modern work on the theory of atomic structure and its bearing on the periodic classification and the positions of the indio-elements is given in this series, Vol I pt. 1, pp. 276-282. For a fuller account, see Soddy, opus cit

 Boltwood, Phys. Review, 1906, 22, 320, Aner. J. Sci., 1906, [iv.], 22, 587; 1908, [iv.], 25, 269, WCoy, Phil. Mag., 1906, [vi.], II, 177; M'Coy and Ross, J. Amer. Chem. Soc., 1907, 29, 1698.

derived from uranium. The a-ray activity of the actimum products is, however, very small in comparison with that of the uranium-radium series of products in the same nuneral (p. 141). Rutherford accordingly suggested that some member of the manium radium series undergoes disintegration in two ways, so that from that member onwards the manum disintegration series branches into two such series, one of which contains actinium and its products. It would suffice to account for the observed results if about 8 per cent, of the disintegrated uranium followed the actinium branch series 2

Passing over the earlier suggestions as to the precise point at which this branching of the uranium series occurs,3 and viewing the problem in the light of modern knowledge, it is seen that if formed by an a ray change, actinium is derived from a homologue of tantalum, and if formed by a  $\beta$  ray change, it is derived from radium or an isotope of that element ! The production of actimum from radium has been disproved. No isotope of radium derived from uranium is known. Such an element would exist it ionium broke down in two ways and a-rays were emitted in each. However, no actinium could be detected in a concentrated ionium preparation four years after it had been made 6 So far as a radioactive homologue of tantalum is concerned, the element brevium or uranium-\", is the only one known, but its short life and absence of a-radiation completely negative the view that it is the parent of actinium. If, then, actinium is formed in an a-ray change, there is an isotope of brevium still to be discovered. It has been sought in pitchblende, with negative results, but there is no certainty that the chemical methods used would have isolated it <sup>7</sup>. The missing element might arise from mamium X<sub>1</sub><sup>8</sup> or ionium by a  $\beta$ -ray change, but the latter is unlikely in view of the failure to produce actinium from ionium. Another alternative has been proposed, and meets with most favour at the present time (1916). Urumini I or uranium II breaks down in two ways, each of the a ray type, producing manum  $X_1$  or ionium on the one hand, and manum-Y on the other Uranium-Y, by a  $\beta$ -ray change, passes into the missing homologue of tantalium The existence of uranium-Y was announced by Antonov in 1911, and is now fairly well established 9 Assuming the branching to start from manning H, the manium series will then be as follows



Rutherford, Radioactive Transformations (Constable, 1906), p. 177
 Rutherford, Radioactive Substances and their Radiodiens (Cambridge Univ Press,

<sup>&</sup>lt;sup>2</sup> Rutherford, Radioactive Substances and their Radioticus (Cambridge Univ Press, 1912), p 523.

<sup>3</sup> M'Coy and Ross, loc cit., Soddy, Phil Mag., 1909, [vi.], 18, 739., cf. Nicholson, Nature, 1911, 87, 515.

<sup>4</sup> Soddy, Chem. Nous, 1913, 107, 97., Jahrb. Radioakt-v Elektronik, 1913, 10, 188; Fajans, Le Radium, 1913, 10, 61, 171., Her., 1913, 46, 422.

<sup>5</sup> Soddy, Nature, 1913, 91, 634.

<sup>6</sup> Gohring, Physikal Zeitsch. 1914, 15, 642.

<sup>7</sup> Gohring, loc cit.

<sup>8</sup> Hahn and Mettner, Physikal Zeitsch., 1913, 14, 752.

<sup>9</sup> Antonov, Phil Mag., 1911. [vi.], 22, 419., 1913, [vi.], 26, 1058., Fleck., ibid., 1913, [vi.], 25, 710., Soddy, ibid., 1914. [vi.], 27, 215; Hahn and Meitner, Physikal. Zeitsch., 1914, 15, 236.

There is reason to believe that the branching proceeds from uranium-II rather than from uranium I. The two alternatives lead to the respective values 226 and 230 for the atomic weight of actinium. Now for a group of isotopic elements it is the rule that for a ray giving members the stability increases with increase in atomic weight, while for  $\beta$ -ray giving members the reverse is true. The actimum series falls in with this rule (to which, however, there are one or two exceptions) much better if Ac = 226 than if  $Ac - 230^{1}$ 

Atomic Weights of Actinium and its Products,-Until pure actinium salts are prepared the chemical equivalent of actinium cannot be directly measured, and until more accurate measurements of the molecular weight of actinium enanation are obtained, the accurate atomic weight of actinium cannot be calculated from that of the emanation If actinium arises from uranium-II, as has been supposed, it should clearly have the same atomic weight as radium, viz 2260 (or 2262 if calculated from U = 2382 and He = 4.00), on the other hand, if it arises from uranium-1, the atomic weight should be 230 0 (or 230 2) Assuming the origin of actinium from uranium-II, the atomic weights in the actinium series will be as follows -

. Ac RaAc AcX AcEm AcA AcB AcC AcC AcD Atomic weight 226 - 226222218 214210 210 210 206

End Products in the Actinium Series.—The nature of the product arising from AcC' by the expulsion of a-rays is quite unknown, but it should be an isotope of lead. This product, however, only arises from 0.2 per cent of the parent actinium, the remainder of which ultimately passes vid AcC by an a-ray change into AcD, thence by a  $\beta$  ray change into another isotope of lead. The nature of this product, AcE say, is also quite unknown, since it emits no detectable radiation. It is of great interest to determine whether AcE is stable or not - Its most probable atomic weight, as has already been explained, is 206 or 206 2, say 206 1, the next alternative being 210 1.

Now the ultimate end product of uranium through the radium series is a stable isotope of lead of calculated atomic weight 2062 (from U=2382) or 2061 (from Ra=2260), say 2061. The atomic weight of "lead" ex-, tracted from uranium minerals practically free from thorium is always less than 207.2, the atomic weight of lead derived from non-radioactive sources,2 and in the cases of (i) a crystalline specimen of uranimite from Africa and (ii.) a crystalline sample of broggerite from Norway, Honigschinid and Mile. St. Horovitz have found Pb = 206 01 and 206 06 respectively, ie the value for the end product of the uranium-radium series. But if about 8 per cent. of this "lead" came from the actimum side-branch, and AcE = 2101, the "lead" should have an "atomic weight" of 206 42. Hence, if AcE = 210.1, it is an unstable element, and disintegrates while the end product of the

Richards and Lembert, J. Amer. Chem. Soc., 1914, 36, 1329, Maurice Curie, Compt. id., 1914, 158, 1676; Homgschmid and Mile. St. Horovitz, ibid., 1914, 158, 1796

3 Hongschmid and Mile. St. Horovitz, Monatsh., 1915, 36, 355.

<sup>1</sup> Fajans, Le Radium, 1918, 10, 171; Physikal Zeitsch, 1913, 14, 951; cf., however, Richardson, Phil May, 1911, [v1], 27, 252. According to S. Meyer, Hess, and Paneth (Sitzungsber. K. Akad Wiss Wica, 1914, 123, 11. a, 1459), when the logarithms of the radioactive constants (A) are plotted against the logarithms of the ranges of the a-rays at N.T.P. the points he on a straight line which cuts the corresponding line for the manum series at UII, indicating the genesis of actinium from that element.

uranium-radium series accumulates. If, on the other hand,  $AcE=206\cdot 1$ , no conclusion as to its stability can be drawn from this line of argument.

Should AcE slowly decay and emit \$\beta\$ rays it would pass into an isotope of bismuth, and if this were stable, the atomic weight of bismuth extracted from uranium minerals would not be 208, but would approach, probably, either 206 or 210. Thus is a matter that is deserving of experimental study. Detection and Estimation of Actinium—For these topics the reader

must be referred to the literature 2

Summary -The properties of actinum and its disintegration products are summarised in the accompanying table

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